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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 7:		(11) International Publication Number: WO 00/24782
C07K 19/00, C12N 15/62, 15/70, 1/21	A2	(43) International Publication Date: 4 May 2000 (04.05.00)
(21) International Application Number: PCT/US (22) International Filing Date: 25 October 1999 (2) (30) Priority Data: 60/105,371 23 October 1998 (23.10.98) 09/428,082 22 October 1999 (22.10.99) (71) Applicant: AMGEN INC. [US/US]; One Amgen Cent Thousand Oaks, CA 91320–1799 (US).	25.10.9 (BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS MW, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC
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(54) Title: MODIFIED PEPTIDES AS THERAPEUTIC AGENTS

(57) Abstract

The present invention concerns fusion of Fc domains with biologically active peptides and a process for preparing pharmaceutical agents using biologically active peptides. In this invention, pharmacologically active compounds are prepared by a process comprising: a) selecting at least one peptide that modulates the activity of a protein of interest; and b) preparing a pharmacologic agent comprising an Fc domain covalently linked to at least one amino acid of the selected peptide. Linkage to the vehicle increases the half-life of the peptide, which otherwise would be quickly degraded in vivo. The preferred vehicle is an Fc domain. The peptide is preferably selected by phage display, E. coli display, ribosome display, RNA-peptide screening, or chemical-peptide screening.

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Modified Peptides as Therapeutic Agents Background of the Invention

Recombinant proteins are an emerging class of therapeutic agents.

Such recombinant therapeutics have engendered advances in protein formulation and chemical modification. Such modifications can protect therapeutic proteins, primarily by blocking their exposure to proteolytic enzymes. Protein modifications may also increase the therapeutic protein's stability, circulation time, and biological activity. A review article describing protein modification and fusion proteins is Francis (1992), Focus on Growth Factors 3:4-10 (Mediscript, London), which is hereby incorporated by reference.

One useful modification is combination with the "Fc" domain of an antibody. Antibodies comprise two functionally independent parts, a variable domain known as "Fab", which binds antigen, and a constant domain known as "Fc", which links to such effector functions as complement activation and attack by phagocytic cells. An Fc has a long serum half-life, whereas an Fab is short-lived. Capon et al. (1989), Nature 337: 525-31. When constructed together with a therapeutic protein, an Fc domain can provide longer half-life or incorporate such functions as Fc receptor binding, protein A binding, complement fixation and perhaps even placental transfer. Id. Table 1 summarizes use of Fc fusions known in the art.

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Table 1—Fc fusion with therapeutic proteins

Form of Fc	Fusion partner	Therapeutic implications	Reference
lgG1	N-terminus of CD30-L	Hodgkin's disease; anaplastic lymphoma; T- cell leukemia	U.S. Patent No. 5,480,981
Murine Fcy2a	IL-10	anti-inflammatory; transplant rejection	Zheng <u>et al</u> . (1995), <u>J.</u> <u>Immunol</u> . 154: 5590-600
lgG1	TNF receptor	septic shock	Fisher <u>et al.</u> (1996), <u>N. Engl. J. Med.</u> 334: 1697-1702; Van Zee, K. <u>et al.</u> (1996), <u>J. Immunol.</u> 156: 2221-30
IgG, IgA, IgM, or IgE (excluding the first domain)	TNF receptor	inflammation, autoimmune disorders	U.S. Pat. No. 5,808,029, issued September 15, 1998
lgG1	CD4 receptor	AIDS	Capon <u>et al.</u> (1989), <u>Nature 337</u> : 525-31
IgG1, IgG3	N-terminus of IL-2	anti-cancer, antiviral	Harvill <u>et al.</u> (1995), <u>Immunotech</u> . 1: 95-105
lgG1	C-terminus of OPG	osteoarthritis; bone density	WO 97/23614, published July 3, 1997
lgG1	N-terminus of leptin	anti-obesity	PCT/US 97/23183, filed December 11, 1997
Human Ig Cγ1	CTLA-4	autoimmune disorders	Linsley (1991), <u>J. Exp.</u> <u>Med</u> . 174:561-9

A much different approach to development of therapeutic agents is peptide library screening. The interaction of a protein ligand with its receptor often takes place at a relatively large interface. However, as demonstrated for human growth hormone and its receptor, only a few key residues at the interface contribute to most of the binding energy. Clackson et al. (1995), Science 267: 383-6. The bulk of the protein ligand merely displays the binding epitopes in the right topology or serves functions unrelated to binding. Thus, molecules of only "peptide" length (2 to 40 amino acids) can bind to the receptor protein of a given large protein ligand. Such peptides may mimic the bioactivity of the large protein ligand ("peptide agonists") or, through competitive binding, inhibit the bioactivity of the large protein ligand ("peptide antagonists").

Phage display peptide libraries have emerged as a powerful method in identifying such peptide agonists and antagonists. See, for example, Scott et al. (1990), Science 249: 386; Devlin et al. (1990), Science 249: 404; U.S. Pat. No. 5,223,409, issued June 29, 1993; U.S. Pat. No. 5,733,731, issued March 31, 1998; U.S. Pat. No. 5,498,530, issued March 12, 5 1996; U.S. Pat. No. 5,432,018, issued July 11, 1995; U.S. Pat. No. 5,338,665, issued August 16, 1994; U.S. Pat. No. 5,922,545, issued July 13, 1999; WO 96/40987, published December 19, 1996; and WO 98/15833, published April 16, 1998 (each of which is incorporated by reference). In such libraries, random peptide sequences are displayed by fusion with coat 10 proteins of filamentous phage. Typically, the displayed peptides are affinity-eluted against an antibody-immobilized extracellular domain of a receptor. The retained phages may be enriched by successive rounds of affinity purification and repropagation. The best binding peptides may be sequenced to identify key residues within one or more structurally related 15 families of peptides. See, e.g., Cwirla et al. (1997), Science 276: 1696-9, in which two distinct families were identified. The peptide sequences may also suggest which residues may be safely replaced by alanine scanning or by mutagenesis at the DNA level. Mutagenesis libraries may be created and screened to further optimize the sequence of the best binders. 20 Lowman (1997), Ann. Rev. Biophys. Biomol. Struct. 26: 401-24.

Structural analysis of protein-protein interaction may also be used to suggest peptides that mimic the binding activity of large protein ligands. In such an analysis, the crystal structure may suggest the identity and relative orientation of critical residues of the large protein ligand, from which a peptide may be designed. See, e.g., Takasaki et al. (1997), Nature Biotech. 15: 1266-70. These analytical methods may also be used to investigate the interaction between a receptor protein and peptides

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selected by phage display, which may suggest further modification of the peptides to increase binding affinity.

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Other methods compete with phage display in peptide research. A peptide library can be fused to the carboxyl terminus of the lac repressor and expressed in E. coli. Another E. coli-based method allows display on the cell's outer membrane by fusion with a peptidoglycan-associated lipoprotein (PAL). Hereinafter, these and related methods are collectively referred to as "E. coli display." In another method, translation of random RNA is halted prior to ribosome release, resulting in a library of polypeptides with their associated RNA still attached. Hereinafter, this and related methods are collectively referred to as "ribosome display." Other methods employ chemical linkage of peptides to RNA; see, for example, Roberts & Szostak (1997), Proc. Natl. Acad. Sci. USA, 94: 12297-303. Hereinafter, this and related methods are collectively referred to as "RNA-peptide screening." Chemically derived peptide libraries have been developed in which peptides are immobilized on stable, non-biological materials, such as polyethylene rods or solvent-permeable resins. Another chemically derived peptide library uses photolithography to scan peptides immobilized on glass slides. Hereinafter, these and related methods are collectively referred to as "chemical-peptide screening." Chemical-peptide screening may be advantageous in that it allows use of D-amino acids and other unnatural analogues, as well as non-peptide elements. Both biological and chemical methods are reviewed in Wells & Lowman (1992), Curr. Opin. Biotechnol. 3: 355-62.

Conceptually, one may discover peptide mimetics of any protein using phage display and the other methods mentioned above. These methods have been used for epitope mapping, for identification of critical amino acids in protein-protein interactions, and as leads for the discovery of new therapeutic agents. E.g., Cortese et al. (1996), Curr. Opin. Biotech. 7:

616-21. Peptide libraries are now being used most often in immunological studies, such as epitope mapping. Kreeger (1996), <u>The Scientist</u> 10(13): 19-20.

Of particular interest here is use of peptide libraries and other techniques in the discovery of pharmacologically active peptides. A number of such peptides identified in the art are summarized in Table 2. The peptides are described in the listed publications, each of which is hereby incorporated by reference. The pharmacologic activity of the peptides is described, and in many instances is followed by a shorthand term therefor in parentheses. Some of these peptides have been modified (e.g., to form C-terminally cross-linked dimers). Typically, peptide libraries were screened for binding to a receptor for a pharmacologically active protein (e.g., EPO receptor). In at least one instance (CTLA4), the peptide library was screened for binding to a monclonal antibody.

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Table 2—Pharmacologically active peptides

Form of peptide	Binding partner/ protein of interest	Pharmacologic activity	Reference
intrapeptide disulfide- bonded	EPO receptor	EPO-mimetic	Wrighton et al. (1996), Science 273: 458-63; U.S. Pat. No. 5,773,569, issued June 30, 1998 to Wrighton et al.
C-terminally cross-linked dimer	EPO receptor	EPO-mimetic	Livnah et al. (1996), Science 273: 464-71; Wrighton et al. (1997), Nature Biotechnology 15: 1261-5; International patent application WO 96/40772, published Dec. 19, 1996
linear	EPO receptor	EPO-mimetic	Naranda <u>et al</u> . (1999), <u>Proc. Natl. Acad. Sci.</u> <u>USA</u> , 96: 7569-74
linear	c-MpI	TPO-mimetic	Cwirla et al. (1997) Science 276: 1696-9; U.S. Pat. No. 5,869,451, issued Feb. 9, 1999; U.S Pat. No. 5,932,946, issued Aug. 3, 1999
C-terminally cross-linked dimer	с-МрІ	TPO-mimetic	Cwirla <u>et al</u> . (1997), <u>Science</u> 276: 1696-9
disulfide- linked dimer		stimulation of hematopoiesis ("G-CSF-mimetic")	Paukovits <u>et al</u> . (1984), <u>Hoppe-Seylers Z.</u> <u>Physiol. Chem</u> . 365: 303 11; Laerum <u>et al</u> . (1988) <u>Exp. Hemat</u> . 16: 274-80
alkylene- linked dimer		G-CSF-mimetic	Bhatnagar <u>et al.</u> (1996), <u>J. Med. Chem.</u> 39: 3814 9; Cuthbertson <u>et al.</u> (1997), <u>J. Med. Chem.</u> 40: 2876-82; King <u>et al.</u> (1991), <u>Exp. Hematol.</u> 19:481; King <u>et al.</u> (1995), <u>Blood.</u> 86 (Suppl.): 309a
linear	IL-1 receptor	inflammatory and autoimmune diseases ("IL-1 antagonist" or "IL-1 ra-mimetic")	U.S. Pat. No. 5,608,035 U.S. Pat. No. 5,786,331 U.S. Pat. No. 5,880,096 Yanofsky et al. (1996),

^{*}The protein listed in this column may be bound by the associated peptide (e.g., EPO receptor, IL-1 receptor) or mimicked by the associated peptide. The references listed for each clarify whether the molecule is bound by or mimicked by the peptides.

		·	Proc. Natl. Acad. Sci. 93: 7381-6; Akeson et al. (1996), J. Biol. Chem. 271: 30517-23; Wiekzorek et al. (1997), Pol. J. Pharmacol. 49: 107-17; Yanofsky (1996), PNAs, 93:7381-7386.
linear	Facteur	stimulation of lymphocytes	inagaki-Ohara <u>et al</u> . (1996), <u>Cellular Immunol</u> .
	thymique serique (FTS)	("FTS-mimetic")	171: 30-40; Yoshida
	Sonquo (1 10)	((1984), l <u>nt. J.</u>
			Immunopharmacol,
introportido	CTLA4 MAb	CTLA4-mimetic	6:141-6. Fukumoto et al. (1998),
intrapeptide disulfide	CILA4 IVIAD	O I EA4-IIIIII duo	Nature Biotech, 16: 267-
bonded			70
exocyclic	TNF-α receptor	TNF-α antagonist	Takasaki <u>et al</u> . (1997), <u>Nature Biotech</u> . 15:1266- 70; WO 98/53842, published December 3, 1998
linear	TNF-α receptor	TNF-α antagonist	Chirinos-Rojas (), <u>J.</u> Imm., 5621-5626.
intrapeptide	C3b	inhibition of complement	Sahu <u>et al</u> . (1996), <u>J.</u>
disulfide		activation; autoimmune	Immunol. 157: 884-91;
bonded		diseases	Morikis <u>et al</u> . (1998), <u>Protein Sci</u> . 7: 619-27
linear	vinculin	("C3b-antagonist") cell adhesion processes—	Adey et al. (1997),
mear	VIIICUIII	cell growth, differentiation,	Biochem. J. 324: 523-8
		wound healing, tumor	
		metastasis ("vinculin	,
linear	C4 binding	binding") anti-thrombotic	Linse et al. (1997), J.
IIIIBAI	protein (C4BP)		Biol. Chem. 272: 14658-
	,		65
linear	urokinase	processes associated with	Goodson <u>et al</u> . (1994), Proc. Natl. Acad. Sci. 91:
	receptor	urokinase interaction with its receptor (e.g.,	7129-33; International
		angiogenesis, tumor cell	application WO
		invasion and metastasis);	97/35969, published
		("UKR antagonist")	October 2, 1997
linear	Mdm2, Hdm2	Inhibition of inactivation of	Picksley et al. (1994),
		p53 mediated by Mdm2 or	Oncogene 9: 2523-9;
		hdm2; anti-tumor ("Mdm/hdm antagonist")	Bottger <u>et al</u> . (1997) <u>J.</u> <u>Mol. Biol</u> . 269: 744-56;
		(Manyham antagonist)	Bottger et al. (1996),
			Oncogene 13: 2141-7
··· linear	p21 ^{WAF1}	anti-tumor by mimicking	-Ball et al. (1997), Curr.
		the activity of p21 ^{waf1}	Biol. 7: 71-80
linear	farnesyl	anti-cancer by preventing	Gibbs et al. (1994), Cell

^b FTS is a thymic hormone mimicked by the molecule of this invention rather than a receptor bound by the molecule of this invention.

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	transferase		77:175-178
linear	Ras effector domain	anti-cancer by inhibiting biological function of the ras oncogene	Moodie et al. (1994), <u>Trends Genet</u> 10: 44-48 Rodriguez et al. (1994), <u>Nature</u> 370:527-532
linear	SH2/SH3 domains	tumor growth with activated tyrosine kinases	Pawson et al (1993), <u>Curr. Biol.</u> 3:434-432 Yu et al. (1994), <u>Cell</u> 76:933-945
linear	p16 ^{iNK4}	anti-cancer by mimicking activity of p16; e.g., inhibiting cyclin D-Cdk complex ("p16-mimetic")	Fáhraeus <u>et al</u> . (1996), <u>Curr. Biol</u> . 6:84-91
linear	Src, Lyn	inhibition of Mast cell activation, IgE-related conditions, type I hypersensitivity ("Mast cell antagonist")	Stauffer et al. (1997), Biochem. 36: 9388-94
linear	Mast cell protease	treatment of inflammatory disorders mediated by release of tryptase-6 ("Mast cell protease inhibitors")	International application WO 98/33812, published August 6, 1998
linear	SH3 domains	treatment of SH3- mediated disease states ("SH3 antagonist")	Rickles et al. (1994), <u>EMBO J</u> . 13: 5598-5604; Sparks et al. (1994), <u>J</u> . <u>Biol. Chem</u> . 269: 23853- 6; Sparks et al. (1996), <u>Proc. Natl. Acad. Sci</u> . 93: 1540-4
linear	HBV core antigen (HBcAg)	treatment of HBV viral infections ("anti-HBV")	Dyson & Muray (1995), <u>Proc. Natl. Acad. Sci</u> . 92: 2194-8
linear	selectins	neutrophil adhesion; inflammatory diseases ("selectin antagonist")	Martens et al. (1995), J. Biol. Chem. 270: 21129-36; European patent application EP 0 714 912, published June 5, 1996
linear, cyclized	calmodulin	calmodulin antagonist	Pierce et al. (1995), Molec. Diversity 1: 259- 65; Dedman et al. (1993), J. Biol. Chem. 268: 23025-30; Adey & Kay (1996), Gene 169: 133-4 International applications
linear, cyclized-	integrins	tumor-homing; treatment for conditions related to integrin-mediated cellular events, including platelet aggregation, thrombosis, wound healing, osteoporosis, tissue repair, angiogenesis (e.g.	WO 95/14714, published June 1, 1995; WO 97/08203, published March 6, 1997; WO 98/10795, published March 19, 1998; WO

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		for treatment of cancer), and tumor invasion ("integrin-binding")	20, 1999; Kraft <u>et al</u> . (1999), J. Biol. Chem. 274: 1979-1985
			WO 98/09985, published
cyclic, linear	fibronectin and extracellular matrix	treatment of inflammatory and autoimmune conditions	March 12, 1998
	components of T cells and macrophages		
lingar	somatostatin	treatment or prevention of	European patent
linear	and cortistatin	hormone-producing tumors, acromegaly, giantism, dementia, gastric ulcer, tumor	application 0 911 393, published April 28, 1999
•		growth, inhibition of hormone secretion, modulation of sleep or neural activity	
		antibiotic; septic shock;	U.S. Pat. No. 5,877,151,
linear	bacterial lipopolysac- charide	disorders modulatable by CAP37	issued March 2, 1999
linear or	pardaxin, mellitin	antipathogenic	WO 97/31019, published
cyclic, including D-	paraaxiii, memiii	• -	28 August 1997
amino acids			WO 97/40070, published
linear, cyclic	VIP	impotence, neurodegenerative disorders	October 30, 1997
linear	CTLs	cancer	EP 0 770 624, published May 2, 1997
linear	THF-gamma2		Burnstein (1988), Biochem., 27:4066-71.
linear	Amylin		Cooper (1987), Proc. Natl. Acad. Sci., 84:8628-32.
linear	Adrenomedullin		Kitamura (1993), <u>BBRC</u> , 192:553-60.
cyclic, linear	VEGF	anti-angiogenic; cancer, rheumatoid arthritis, diabetic retinopathy, psoriasis ("VEGF antagonist")	Fairbrother (1998), <u>Biochem.,</u> 37:17754- 17764.
cyclic	ММР	inflammation and autoimmune disorders; tumor growth ("MMP inhibitor")	Koivunen (1999), <u>Nature</u> <u>Biotech.</u> , 17:768-774.
	HGH fragment		U.S. Pat. No. 5,869,452
	Echistatin	inhibition of platelet aggregation	Gan (1988), <u>J. Biol.</u> Chem., 263:19827-32.
linear	SLE autoantibody	SLE	WO 96/30057, publishe October 3, 1996 Ishikawa et al. (1998),
	GD1alpha	suppression of tumor metastasis	FEBS Lett. 441 (1): 20-
	antiphospholipio	endothelial cell activation	, Blank et al. (1999), Proc

	beta-2- glycoprotein-l (β2GPI) antibodies	antiphospholipid syndrome (APS), thromboembolic phenomena, thrombocytopenia, and recurrent fetal loss	Natl. Acad. Sci. USA 96: 5164-8
linear	T Cell Receptor beta chain	diabetes	WO 96/11214, published April 18, 1996

Peptides identified by peptide library screening have been regarded as "leads" in development of therapeutic agents rather than as therapeutic agents themselves. Like other proteins and peptides, they would be rapidly removed in vivo either by renal filtration, cellular clearance mechanisms in the reticuloendothelial system, or proteolytic degradation. Francis (1992), Focus on Growth Factors 3: 4-11. As a result, the art presently uses the identified peptides to validate drug targets or as scaffolds for design of organic compounds that might not have been as easily or as quickly identified through chemical library screening. Lowman (1997), Ann. Rev. Biophys. Biomol. Struct. 26: 401-24; Kay et al. (1998), Drug Disc. Today 3: 370-8. The art would benefit from a process by which such peptides could more readily yield therapeutic agents.

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Summary of the Invention

The present invention concerns a process by which the <u>in vivo</u> halflife of one or more biologically active peptides is increased by fusion with a vehicle. In this invention, pharmacologically active compounds are prepared by a process comprising:

- a) selecting at least one peptide that modulates the activity of a protein of interest; and
- b) preparing a pharmacologic agent comprising at least one vehicle covalently linked to at least one amino acid sequence of the selected peptide.

The preferred vehicle is an Fc domain. The peptides screened in step (a) are preferably expressed in a phage display library. The vehicle and the

peptide may be linked through the N- or C-terminus of the peptide or the vehicle, as described further below. Derivatives of the above compounds (described below) are also encompassed by this invention.

The compounds of this invention may be prepared by standard synthetic methods, recombinant DNA techniques, or any other methods of preparing peptides and fusion proteins. Compounds of this invention that encompass non-peptide portions may be synthesized by standard organic chemistry reactions, in addition to standard peptide chemistry reactions when applicable.

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The primary use contemplated is as therapeutic or prophylactic agents. The vehicle-linked peptide may have activity comparable to—or even greater than—the natural ligand mimicked by the peptide. In addition, certain natural ligand-based therapeutic agents might induce antibodies against the patient's own endogenous ligand; the vehicle-linked peptide avoids this pitfall by having little or typically no sequence identity with the natural ligand.

Although mostly contemplated as therapeutic agents, compounds of this invention may also be useful in screening for such agents. For example, one could use an Fc-peptide (e.g., Fc-SH2 domain peptide) in an assay employing anti-Fc coated plates. The vehicle, especially Fc, may make insoluble peptides soluble and thus useful in a number of assays.

The compounds of this invention may be used for therapeutic or prophylactic purposes by formulating them with appropriate pharmaceutical carrier materials and administering an effective amount to a patient, such as a human (or other mammal) in need thereof. Other related aspects are also included in the instant invention.

Numerous additional aspects and advantages of the present invention will become apparent upon consideration of the figures and detailed description of the invention.

Brief Description of the Figures

Figure 1 shows a schematic representation of an exemplary process of the invention. In this preferred process, the vehicle is an Fc domain, which is linked to the peptide covalently by expression from a DNA construct encoding both the Fc domain and the peptide. As noted in Figure 1, the Fc domains spontaneously form a dimer in this process.

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Figure 2 shows exemplary Fc dimers that may be derived from an IgG1 antibody. "Fc" in the figure represents any of the Fc variants within the meaning of "Fc domain" herein. "X¹" and "X²" represent peptides or linker-peptide combinations as defined hereinafter. The specific dimers are as follows:

A, D: Single disulfide-bonded dimers. IgG1 antibodies typically have two disulfide bonds at the hinge region between the constant and variable domains. The Fc domain in Figures 2A and 2 D may be formed by truncation between the two disulfide bond sites or by substitution of a cysteinyl residue with an unreactive residue (e.g., alanyl). In Figure 2A, the Fc domain is linked at the amino terminus of the peptides; in 2D, at the carboxyl terminus.

B, E: Doubly disulfide-bonded dimers. This Fc domain may be formed by truncation of the parent antibody to retain both cysteinyl residues in the Fc domain chains or by expression from a construct including a sequence encoding such an Fc domain. In Figure 2B, the Fc domain is linked at the amino terminus of the peptides; in 2E, at the carboxyl terminus.

C, F: Noncovalent dimers. This Fc domain may be formed by elimination of the cysteinyl residues by either truncation or substitution.

One may desire to eliminate the cysteinyl residues to avoid impurities formed by reaction of the cysteinyl residue with cysteinyl residues of other

proteins present in the host cell. The noncovalent bonding of the Fc domains is sufficient to hold together the dimer.

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Other dimers may be formed by using Fc domains derived from different types of antibodies (e.g., IgG2, IgM).

Figure 3 shows the structure of preferred compounds of the invention that feature tandem repeats of the pharmacologically active peptide. Figure 3A shows a single chain molecule and may also represent the DNA construct for the molecule. Figure 3B shows a dimer in which the linker-peptide portion is present on only one chain of the dimer. Figure 3C shows a dimer having the peptide portion on both chains. The dimer of Figure 3C will form spontaneously in certain host cells upon expression of a DNA construct encoding the single chain shown in Figure 3A. In other host cells, the cells could be placed in conditions favoring formation of dimers or the dimers can be formed in vitro.

Figure 4 shows exemplary nucleic acid and amino acid sequences (SEQ ID NOS: 1 and 2, respectively) of human IgG1 Fc that may be used in this invention.

Figure 5 shows a synthetic scheme for the preparation of PEGylated peptide 19 (SEQ ID NO: 3).

Figure 6 shows a synthetic scheme for the preparation of PEGylated peptide 20 (SEQ ID NO: 4).

Figure 7 shows the nucleotide and amino acid sequences (SEQ ID NOS: 5 and 6, respectively) of the molecule identified as "Fc-TMP" in Example 2 hereinafter.

Figure 8 shows the nucleotide and amino acid sequences (SEQ. ID. NOS: 7 and 8, respectively) of the molecule identified as "Fc-TMP-TMP" in Example 2 hereinafter.

Figure 9 shows the nucleotide and amino acid sequences (SEQ. ID. NOS: 9 and 10, respectively) of the molecule identified as "TMP-TMP-Fc" in Example 2 hereinafter.

Figure 10 shows the nucleotide and amino acid sequences (SEQ. ID. NOS: 11 and 12, respectively) of the molecule identified as "TMP-Fc" in Example 2 hereinafter.

Figure 11 shows the number of platelets generated <u>in vivo</u> in normal female BDF1 mice treated with one 100 μ g/kg bolus injection of various compounds, with the terms defined as follows.

PEG-MGDF: 20 kD average molecular weight PEG attached by reductive amination to the N-terminal amino group of amino acids 1-163 of native human TPO, which is expressed in <u>E. coli</u> (so that it is not glycosylated);

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- TMP: the TPO-mimetic peptide having the amino acid sequence IEGPTLRQWLAARA (SEQ ID NO: 13);
- TMP-TMP: the TPO-mimetic peptide having the amino acid sequence IEGPTLRQWLAARA-GGGGGGGG-IEGPTLRQWLAARA (SEQ ID NO: 14);
- PEG-TMP-TMP: the peptide of SEQ ID NO: 14, wherein the PEG group is a 5 kD average molecular weight PEG attached as shown in Figure 6;
- Fc-TMP-TMP: the compound of SEQ ID NO: 8 (Figure 8) dimerized with an identical second monomer (i.e., Cys residues 7 and 10 are bound to the corresponding Cys residues in the second monomer to form a dimer, as shown in Figure 2); and
- TMP-TMP-Fc is the compound of SEQ ID NO: 10 (Figure 9)
 dimerized in the same way as TMP-TMP-Fc except that the Fc.
 domain is attached at the C-terminal end rather than the Nterminal end of the TMP-TMP peptide.

Figure 12 shows the number of platelets generated <u>in vivo</u> in normal BDF1 mice treated with various compounds delivered via implanted osmotic pumps over a 7-day period. The compounds are as defined for Figure 7.

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Figure 13 shows the nucleotide and amino acid sequences (SEQ. ID. NOS: 15 and 16, respectively) of the molecule identified as "Fc-EMP" in Example 3 hereinafter.

Figure 14 shows the nucleotide and amino acid sequences (SEQ ID NOS: 17 and 18, respectively) of the molecule identified as "EMP-Fc" in Example 3 hereinafter.

Figure 15 shows the nucleotide and amino acid sequences (SEQ ID NOS:19 and 20, respectively) of the molecule identified as "EMP-EMP-Fc" in Example 3 hereinafter.

Figure 16 shows the nucleotide and amino acid sequences (SEQ ID NOS: 21 and 22, respectively) of the molecule identified as "Fc-EMP-EMP" in Example 3 hereinafter.

Figures 17A and 17B show the DNA sequence (SEQ ID NO: 23) inserted into pCFM1656 between the unique <u>Aat</u>II (position #4364 in pCFM1656) and <u>Sac</u>II (position #4585 in pCFM1656) restriction sites to form expression plasmid pAMG21 (ATCC accession no. 98113).

Figure 18A shows the hemoglobin, red blood cells, and hematocrit generated in vivo in normal female BDF1 mice treated with one 100 μ g/kg bolus injection of various compounds. Figure 18B shows the same results with mice treated with 100 μ g/kg per day delivered the same dose by 7-day micro-osmotic pump with the EMPs delivered at 100 μ g/kg, rhEPO at 30U/mouse. (In both experiments, neutrophils, lymphocytes, and platelets were unaffected.) In these figures, the terms are defined as follows.

Fc-EMP: the compound of SEQ ID NO: 16 (Figure 13) dimerized with an identical second monomer (i.e., Cys residues 7 and 10 are

bound to the corresponding Cys residues in the second monomer to form a dimer, as shown in Figure 2);

EMP-Fc: the compound of SEQ ID NO: 18 (Figure 14) dimerized in the same way as Fc-EMP except that the Fc domain is attached at the C-terminal end rather than the N-terminal end of the EMP peptide.

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"EMP-EMP-Fc" refers to a tandem repeat of the same peptide (SEQ ID NO: 20) attached to the same Fc domain by the carboxyl terminus of the peptides. "Fc-EMP-EMP" refers to the same tandem repeat of the peptide but with the same Fc domain attached at the amino terminus of the tandem repeat. All molecules are expressed in E. coli and so are not glycosylated.

Figures 19A and 19B show the nucleotide and amino acid sequences (SEQ ID NOS: 1055 and 1056) of the Fc-TNF- α inhibitor fusion molecule described in Example 4 hereinafter.

Figures 20A and 20B show the nucleotide and amino acid sequences (SEQ ID NOS: 1057 and 1058) of the TNF- α inhibitor-Fc fusion molecule described in Example 4 hereinafter.

Figures 21A and 21B show the nucleotide and amino acid sequences (SEQ ID NOS: 1059 and 1060) of the Fc-IL-1 antagonist fusion molecule described in Example 5 hereinafter.

Figures 22A and 22B show the nucleotide and amino acid sequences (SEQ ID NOS: 1061 and 1062) of the IL-1 antagonist-Fc fusion molecule described in Example 5 hereinafter.

Figures 23A, 23B, and 23C show the nucleotide and amino acid sequences (SEQ ID NOS: 1063 and 1064) of the Fc-VEGF antagonist fusion molecule described in Example 6 hereinafter.

Figures 24A and 24B show the nucleotide and amino acid sequences (SEQ ID NOS: 1065 and 1066) of the VEGF antagonist-Fc fusion molecule described in Example 6 hereinafter.

Figures 25A and 25B show the nucleotide and amino acid sequences (SEQ ID NOS: 1067 and 1068) of the Fc-MMP inhibitor fusion molecule described in Example 7 hereinafter.

Figures 26A and 26B show the nucleotide and amino acid sequences (SEQ ID NOS: 1069 and 1070) of the MMP inhibitor-Fc fusion molecule described in Example 7 hereinafter.

Detailed Description of the Invention

Definition of Terms

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The terms used throughout this specification are defined as follows, unless otherwise limited in specific instances.

The term "comprising" means that a compound may include additional amino acids on either or both of the N- or C- termini of the given sequence. Of course, these additional amino acids should not significantly interfere with the activity of the compound.

The term "vehicle" refers to a molecule that prevents degradation and/or increases half-life, reduces toxicity, reduces immunogenicity, or increases biological activity of a therapeutic protein. Exemplary vehicles include an Fc domain (which is preferred) as well as a linear polymer (e.g., polyethylene glycol (PEG), polylysine, dextran, etc.); a branched-chain polymer (see, for example, U.S. Patent No. 4,289,872 to Denkenwalter et al., issued September 15, 1981; 5,229,490 to Tam, issued July 20, 1993; WO 93/21259 by Frechet et al., published 28 October 1993); a lipid; a cholesterol group (such as a steroid); a carbohydrate or oligosaccharide; or any natural or synthetic protein, polypeptide or peptide that binds to a salvage receptor. Vehicles are further described hereinafter.

The term "native Fc" refers to molecule or sequence comprising the sequence of a non-antigen-binding fragment resulting from digestion of whole antibody, whether in monomeric or multimeric form. The original immunoglobulin source of the native Fc is preferably of human origin and may be any of the immunoglobulins, although IgG1 and IgG2 are preferred. Native Fc's are made up of monomeric polypeptides that may be linked into dimeric or multimeric forms by covalent (i.e., disulfide bonds) and non-covalent association. The number of intermolecular disulfide bonds between monomeric subunits of native Fc molecules ranges from 1 to 4 depending on class (e.g., IgG, IgA, IgE) or subclass (e.g., IgG1, IgG2, IgG3, IgA1, IgGA2). One example of a native Fc is a disulfide-bonded dimer resulting from papain digestion of an IgG (see Ellison et al. (1982), Nucleic Acids Res. 10: 4071-9). The term "native Fc" as used herein is generic to the monomeric, dimeric, and multimeric forms.

The term "Fc variant" refers to a molecule or sequence that is modified from a native Fc but still comprises a binding site for the salvage receptor, FcRn. International applications WO 97/34631 (published 25 September 1997) and WO 96/32478 describe exemplary Fc variants, as well as interaction with the salvage receptor, and are hereby incorporated by reference. Thus, the term "Fc variant" comprises a molecule or sequence that is humanized from a non-human native Fc. Furthermore, a native Fc comprises sites that may be removed because they provide structural features or biological activity that are not required for the fusion molecules of the present invention. Thus, the term "Fc variant" comprises a molecule or sequence that lacks one or more native Fc sites or residues that affect or are involved in (1) disulfide bond formation, (2) incompatibility with a selected host cell (3) N-terminal heterogeneity upon expression in a selected host cell, (4) glycosylation, (5) interaction with complement, (6) binding to an Fc receptor other than a salvage receptor, or

(7) antibody-dependent cellular cytotoxicity (ADCC). Fc variants are described in further detail hereinafter.

The term "Fc domain" encompasses native Fc and Fc variant molecules and sequences as defined above. As with Fc variants and native Fc's, the term "Fc domain" includes molecules in monomeric or multimeric form, whether digested from whole antibody or produced by other means.

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The term "multimer" as applied to Fc domains or molecules comprising Fc domains refers to molecules having two or more polypeptide chains associated covalently, noncovalently, or by both covalent and non-covalent interactions. IgG molecules typically form dimers; IgM, pentamers; IgD, dimers; and IgA, monomers, dimers, trimers, or tetramers. Multimers may be formed by exploiting the sequence and resulting activity of the native Ig source of the Fc or by derivatizing (as defined below) such a native Fc.

The term "dimer" as applied to Fc domains or molecules comprising Fc domains refers to molecules having two polypeptide chains associated covalently or non-covalently. Thus, exemplary dimers within the scope of this invention are as shown in Figure 2.

The terms "derivatizing" and "derivative" or "derivatized" comprise processes and resulting compounds respectively in which (1) the compound has a cyclic portion; for example, cross-linking between cysteinyl residues within the compound; (2) the compound is cross-linked or has a cross-linking site; for example, the compound has a cysteinyl residue and thus forms cross-linked dimers in culture or in vivo; (3) one or more peptidyl linkage is replaced by a non-peptidyl linkage; (4) the N-terminus is replaced by -NRR¹, NRC(O)R¹, -NRC(O)OR¹, -NRS(O)₂R¹, -NHC(O)NHR, a succinimide group, or substituted or unsubstituted benzyloxycarbonyl-NH-, wherein R and R¹ and the ring substituents are

as defined hereinafter; (5) the C-terminus is replaced by -C(O)R² or -NR³R⁴ wherein R², R³ and R⁴ are as defined hereinafter; and (6) compounds in which individual amino acid moieties are modified through treatment with agents capable of reacting with selected side chains or terminal residues. Derivatives are further described hereinafter.

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The term "peptide" refers to molecules of 2 to 40 amino acids, with molecules of 3 to 20 amino acids preferred and those of 6 to 15 amino acids most preferred. Exemplary peptides may be randomly generated by any of the methods cited above, carried in a peptide library (e.g., a phage display library), or derived by digestion of proteins.

The term "randomized" as used to refer to peptide sequences refers to fully random sequences (e.g., selected by phage display methods) and sequences in which one or more residues of a naturally occurring molecule is replaced by an amino acid residue not appearing in that position in the naturally occurring molecule. Exemplary methods for identifying peptide sequences include phage display, <u>E. coli</u> display, ribosome display, RNA-peptide screening, chemical screening, and the like.

The term "pharmacologically active" means that a substance so described is determined to have activity that affects a medical parameter (e.g., blood pressure, blood cell count, cholesterol level) or disease state (e.g., cancer, autoimmune disorders). Thus, pharmacologically active peptides comprise agonistic or mimetic and antagonistic peptides as defined below.

The terms "-mimetic peptide" and "-agonist peptide" refer to a peptide having biological activity comparable to a protein (e.g., EPO, TPO, G-CSF) that interacts with a protein of interest. These terms further include peptides that indirectly mimic the activity of a protein of interest, such as by potentiating the effects of the natural ligand of the protein of interest; see, for example, the G-CSF-mimetic peptides listed in Tables 2

and 7. Thus, the term "EPO-mimetic peptide" comprises any peptides that can be identified or derived as described in Wrighton et al. (1996), Science 273: 458-63, Naranda et al. (1999), Proc. Natl. Acad. Sci. USA 96: 7569-74, or any other reference in Table 2 identified as having EPO-mimetic subject matter. Those of ordinary skill in the art appreciate that each of these references enables one to select different peptides than actually disclosed therein by following the disclosed procedures with different peptide libraries.

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The term "TPO-mimetic peptide" comprises peptides that can be identified or derived as described in Cwirla et al. (1997), Science 276: 1696-9, U.S. Pat. Nos. 5,869,451 and 5,932,946 and any other reference in Table 2 identifed as having TPO-mimetic subject matter, as well as the U.S. patent application, "Thrombopoietic Compounds," filed on even date herewith and hereby incorporated by reference. Those of ordinary skill in the art appreciate that each of these references enables one to select different peptides than actually disclosed therein by following the disclosed procedures with different peptide libraries.

The term "G-CSF-mimetic peptide" comprises any peptides that can be identified or described in Paukovits et al. (1984), Hoppe-Seylers Z. Physiol. Chem. 365: 303-11 or any of the references in Table 2 identified as having G-CSF-mimetic subject matter. Those of ordinary skill in the art appreciate that each of these references enables one to select different peptides than actually disclosed therein by following the disclosed procedures with different peptide libraries.

The term "CTLA4-mimetic peptide" comprises any peptides that can be identified or derived as described in Fukumoto et al. (1998), Nature Biotech. 16: 267-70. Those of ordinary skill in the art appreciate that each of these references enables one to select different peptides than actually

disclosed therein by following the disclosed procedures with different peptide libraries.

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The term "-antagonist peptide" or "inhibitor peptide" refers to a peptide that blocks or in some way interferes with the biological activity of the associated protein of interest, or has biological activity comparable to a known antagonist or inhibitor of the associated protein of interest. Thus, the term "TNF-antagonist peptide" comprises peptides that can be identified or derived as described in Takasaki et al. (1997), Nature Biotech. 15: 1266-70 or any of the references in Table 2 identified as having TNF-antagonistic subject matter. Those of ordinary skill in the art appreciate that each of these references enables one to select different peptides than actually disclosed therein by following the disclosed procedures with different peptide libraries.

The terms "IL-1 antagonist" and "IL-1ra-mimetic peptide" comprises peptides that inhibit or down-regulate activation of the IL-1 receptor by IL-1. IL-1 receptor activation results from formation of a complex among IL-1, IL-1 receptor, and IL-1 receptor accessory protein. IL-1 antagonist or IL-1ra-mimetic peptides bind to IL-1, IL-1 receptor, or IL-1 receptor accessory protein and obstruct complex formation among any two or three components of the complex. Exemplary IL-1 antagonist or IL-1ra-mimetic peptides can be identified or derived as described in U.S. Pat. Nos. 5,608,035, 5,786,331, 5,880,096, or any of the references in Table 2 identified as having IL-1ra-mimetic or IL-1 antagonistic subject matter. Those of ordinary skill in the art appreciate that each of these references enables one to select different peptides than actually disclosed therein by following the disclosed procedures with different peptide libraries.

The term "VEGF-antagonist peptide" comprises peptides that can be identified or derived as described in Fairbrother (1998), <u>Biochem.</u> 37:

17754-64, and in any of the references in Table 2 identified as having VEGF-antagonistic subject matter. Those of ordinary skill in the art appreciate that each of these references enables one to select different peptides than actually disclosed therein by following the disclosed procedures with different peptide libraries.

The term "MMP inhibitor peptide" comprises peptides that can be identified or derived as described in Koivunen (1999), Nature Biotech. 17: 768-74 and in any of the references in Table 2 identified as having MMP inhibitory subject matter. Those of ordinary skill in the art appreciate that each of these references enables one to select different peptides than actually disclosed therein by following the disclosed procedures with different peptide libraries.

Additionally, physiologically acceptable salts of the compounds of this invention are also encompassed herein. By "physiologically acceptable salts" is meant any salts that are known or later discovered to be pharmaceutically acceptable. Some specific examples are: acetate; trifluoroacetate; hydrohalides, such as hydrochloride and hydrobromide; sulfate; citrate; tartrate; glycolate; and oxalate.

Structure of compounds

In General. In the compositions of matter prepared in accordance with this invention, the peptide may be attached to the vehicle through the peptide's N-terminus or C-terminus. Thus, the vehicle-peptide molecules of this invention may be described by the following formula I:

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$$(X^1)_a - F^1 - (X^2)_b$$

wherein:

F¹ is a vehicle (preferably an Fc domain);

 $X^{1} \text{ and } X^{2} \text{ are each independently selected from -(L^{1})}_{c} - P^{1}, -(L^{1})_{c} - P^{1} - (L^{2})_{d} - P^{2}, -(L^{1})_{c} - P^{1} - (L^{2})_{d} - P^{2} - (L^{3})_{e} - P^{3}, \text{ and -(L^{1})}_{c} - P^{1} - (L^{2})_{d} - P^{2} - (L^{3})_{e} - P^{3} - (L^{4})_{f} - P^{4}$

P¹, P², P³, and P⁴ are each independently sequences of pharmacologically active peptides;

L1, L2, L3, and L4 are each independently linkers; and

a, b, c, d, e, and f are each independently 0 or 1, provided that at least one of a and b is 1.

Thus, compound I comprises preferred compounds of the formulae

X1-F1

and multimers thereof wherein F¹ is an Fc domain and is attached at the Cterminus of X¹;

Ш

П

$$F^1-X^2$$

and multimers thereof wherein F^1 is an Fc domain and is attached at the N-terminus of X^2 ;

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and multimers thereof wherein F^1 is an Fc domain and is attached at the N-terminus of $-(L^1)$, $-P^1$; and

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$$F^1-(L^1)_c-P^1-(L^2)_d-P^2$$

and multimers thereof wherein F^1 is an Fc domain and is attached at the N-terminus of $-L^1-P^1-L^2-P^2$.

<u>Peptides</u>. Any number of peptides may be used in conjunction with the present invention. Of particular interest are peptides that mimic the activity of EPO, TPO, growth hormone, G-CSF, GM-CSF, IL-1ra, leptin, CTLA4, TRAIL, TGF- α , and TGF- β . Peptide antagonists are also of interest, particularly those antagonistic to the activity of TNF, leptin, any of the interleukins (IL-1, 2, 3, ...), and proteins involved in complement activation (e.g., C3b). Targeting peptides are also of interest, including

tumor-homing peptides, membrane-transporting peptides, and the like.

All of these classes of peptides may be discovered by methods described in the references cited in this specification and other references.

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Phage display, in particular, is useful in generating peptides for use in the present invention. It has been stated that affinity selection from libraries of random peptides can be used to identify peptide ligands for any site of any gene product. Dedman et al. (1993), J. Biol. Chem. 268: 23025-30. Phage display is particularly well suited for identifying peptides that bind to such proteins of interest as cell surface receptors or any proteins having linear epitopes. Wilson et al. (1998), Can. J. Microbiol. 44: 313-29; Kay et al. (1998), Drug Disc. Today 3: 370-8. Such proteins are extensively reviewed in Herz et al. (1997), J. Receptor & Signal Transduction Res. 17(5): 671-776, which is hereby incorporated by reference. Such proteins of interest are preferred for use in this invention.

A particularly preferred group of peptides are those that bind to cytokine receptors. Cytokines have recently been classified according to their receptor code. See Inglot (1997), <u>Archivum Immunologiae et Therapiae Experimentalis</u> 45: 353-7, which is hereby incorporated by reference. Among these receptors, most preferred are the CKRs (family I in Table 3). The receptor classification appears in Table 3.

PCT/US99/25044

Table 3—Cytokine Receptors Classified by Receptor Code

Cytokine	s (ligands)	Recept	tor Type
family	subfamily	family	subfamily
I. Hematopoietic cytokines	1. IL-2, IL-4, IL-7, IL-9, IL-13, IL- 15	I. Cytokine R (CKR)	1. shared γCr
	2. IL-3, IL-5, GM- CSF		2. shared GP 140 βR
	3. IL-6, IL-11, IL- 12, LIF, OSM, CNTF, leptin (OB)		3. 3.shared RP 130
	4. G-CSF, EPO, TPO, PRL, GH		4. "single chain" R
	5. IL-17, HVS-IL- 17		5. other R°
II. IL-10 ligands	IL-10, BCRF-1, HSV-IL-10	II. IL-10 R	
III. Interferons	1. IFN-αl, α2, α4, m, t, IFN-β ^d	III. Interferon R	1. IFNAR
	2. IFN-γ		2. IFNGR
IV. IL-1 ligands	1. IL-1α, IL-1β, IL- 1Ra	IV. IL-1R	
V. TNF ligands	1. TNF-α, TNF-β (LT), FAS1, CD40 L, CD30L, CD27 L	V. NGF/TNF R°	
VI. Chemokines	 α chemokines: IL-8, GRO α, β, γ, IF-10, PF-4, SDF-1 	VI. Chemokine R	1. CXCR
	 β chemokines: MIP1α, MIP1β, MCP-1,2,3,4, RANTES, eotaxin 		2. CCR
	 γ chemokines: lymphotactin 		 3. CR 4. DARC'

⁴ Other IFN type I subtypes remain unassigned. Hematopoletic cytokines, IL-10 ligands and interferons do not possess functional intrinsic protein kinases. The signaling molecules for the cytokines are JAK's, STATs and related non-receptor molecules. IL-14, IL-16 and IL-18 have been cloned but according to the receptor code they remain unassigned.

The Duffy blood group antigen (DARC) is an erythrocyte receptor that can bind several different chemokines. It belongs to the immunoglobulin superfamily but characteristics of its signal transduction events remain unclear.

IL-17R belongs to the CKR family but is not assigned to any of the 4 indicated subjamilies.
 Other IFN type I subtypes remain unassigned. Hematopoietic cytokines, IL-10 ligands and

^{*} TNF receptors use multiple, distinct intracellular molecules for signal transduction including "death domain" of FAS R and 55 kDa TNF-αR that participates in their cytotoxic effects. NGF/TNF R can bind both NGF and related factors as well as TNF ligands. Chemokine receptors are G protein-coupled, seven transmembrane (7TM, serpentine) domain receptors.

PCT/US99/25044 WO 00/24782

VII. Growth factors		VII. RKF	1	TK sub-family
VII. Growth factors	4 4 005 11 005	VII. 1 (1.5)		
4	1.1 SCF, M-CSF,		1.1	igTK III R
	PDGF-AA, AB,			
	BB, FLT-3L,			
	VEGF, SSV-			
	· ·			
	PDGF			
	1.2 FGFα, FGFβ	•		IgTK IV R
	1.3 EGF, TGF-α,		1.3	Cysteine-rich
	VV-F19 (EGF-			TK-I
	like)		4.4	Custoine sich
	1.4 IGF-I, IGF-II,		1.4	Cysteine rich
	Insulin			TK-II
	1.5 NGF, BDNF,		1.5	Cysteine knot
	NT-3, NT-4°			TK V
	•		2	STK subfamily ^h
	2. TGF-β1,β2,β3		2.	STA Subiamily

Exemplary peptides for this invention appear in Tables 4 through 20 below. These peptides may be prepared by methods disclosed in the art. Single letter amino acid abbreviations are used. The X in these sequences (and throughout this specification, unless specified otherwise in a particular instance) means that any of the 20 naturally occurring amino acid residues may be present. Any of these peptides may be linked in tandem (i.e., sequentially), with or without linkers, and a few tandemlinked examples are provided in the table. Linkers are listed as " Λ " and 10 may be any of the linkers described herein. Tandem repeats and linkers are shown separated by dashes for clarity. Any peptide containing a cysteinyl residue may be cross-linked with another Cys-containing peptide, either or both of which may be linked to a vehicle. A few crosslinked examples are provided in the table. Any peptide having more than 15 one Cys residue may form an intrapeptide disulfide bond, as well; see, for example, EPO-mimetic peptides in Table 5. A few examples of intrapeptide disulfide-bonded peptides are specified in the table. Any of these peptides may be derivatized as described herein, and a few derivatized examples are provided in the table. Derivatized peptides in 20

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⁹ The neurotrophic cytokines can associate with NGF/TNF receptors also.

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the tables are exemplary rather than limiting, as the associated underivatized peptides may be employed in this invention, as well. For derivatives in which the carboxyl terminus may be capped with an amino group, the capping amino group is shown as -NH₂. For derivatives in which amino acid residues are substituted by moieties other than amino acid residues, the substitutions are denoted by σ , which signifies any of the moieties described in Bhatnagar et al. (1996), J. Med. Chem. 39: 3814-9 and Cuthbertson et al. (1997), J. Med. Chem. 40: 2876-82, which are incorporated by reference. The J substituent and the Z substituents (Z_{z} , Z_{z}) ... Z_{a}) are as defined in U.S. Pat. Nos. 5,608,035,5,786,331, and 5,880,096, which are incorporated by reference. For the EPO-mimetic sequences (Table 5), the substituents X, through X_{11} and the integer "n" are as defined in WO 96/40772, which is incorporated by reference. The substituents "Y," "0," and "+" are as defined in Sparks et al. (1996), Proc. Natl. Acad. Sci. 93: 1540-4, which is hereby incorporated by reference. X_a, X_b, X_b, and X, are as defined in U.S. Pat. No. 5,773,569, which is hereby incorporated by reference, except that: for integrin-binding peptides, X_1 , X_2 , X_3 , X_4 , X_4 , X_5 , X_6 , X_7 , and X₈ are as defined in International applications WO 95/14714, published June 1, 1995 and WO 97/08203, published March 6, 1997, which are also incorporated by reference; and for VIP-mimetic peptides, X,, X,', X_1 ", X_2 , X_3 , X_4 , X_5 , X_6 and Z and the integers m and n are as defined in WO 97/40070, published October 30, 1997, which is also incorporated by reference. Xaa and Yaa below are as defined in WO 98/09985, published March 12, 1998, which is incorporated by reference. AA₁, AA₂, AB₁, AB₂, and AC are as defined in International application WO 98/53842, published December 3, 1998, which is incorporated by reference. X^1 , X^2 , X^3 , and X4 in Table 17 only are as defined in European application EP 0 911

^h STKS may encompass many other TGF-β-related factors that remain unassigned. The protein kinases are intrinsic part of the intracellular domain of receptor kinase family (RKF). The enzymes participate in the signals transmission via the receptors.

393, published April 28, 1999. Residues appearing in boldface are Damino acids. All peptides are linked through peptide bonds unless otherwise noted. Abbreviations are listed at the end of this specification. In the "SEQ ID NO." column, "NR" means that no sequence listing is required for the given sequence.

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Table 4—IL-1 antagonist peptide sequences

Sequence/structure	SEQ
	ID NO:
Z,,Z,Z,QZ,YZ,Z,Q	212
XXQZ,YZ,XX	907
Z,XQZ,YZ,XX	908
Z,Z ₈ QZ ₅ YZ ₆ Z ₃ Z ₁₀	909
Z,ZZ,QZ,YZ,Z,	910
Z ₁₂ Z ₁₃ Z ₁₄ Z ₁₅ Z ₁₆ Z ₁₇ Z ₁₈ Z ₁₃ Z ₂₀ Z ₂₁ Z ₂₂ Z ₁₁ Z ₂ Z ₃ QZ ₅ YZ ₆ Z ₂ Z ₁₀ L	917
Z,NZ,Z,Z,Z,Z,Z,Z,Z,Z,Z,Z,Z,Z,Z,Z,Z,Z,Z,	979
TANVSSFEWTPYYWQPYALPL	213
SWTDYGYWQPYALPISGL	214
ETPFTWEESNAYYWQPYALPL	215
ENTYSPNWADSMYWQPYALPL	216
SVGEDHNFWTSEYWQPYALPL	217
DGYDRWRQSGERYWQPYALPL	218
FEWTPGYWQPY	219
FEWTPGYWQHY	220
FEWTPGWYQJY	221
AcFEWTPGWYQJY	222
FEWTPGWpYQJY	223
FAWTPGYWQJY	224
FEWAPGYWQJY	225
FEWVPGYWQJY	226
FEWTPGYWQJY	227
AcFEWTPGYWQJY	228
FEWTPaWYQJY	229
FEWTPSarWYQJY	230
FEWTPGYYQPY	231
FEWTPGWWQPY	232
FEWTPNYWQPY	233
FEWTPvYWQJY	234
FEWTPecGYWQJY .	235
FEWTPAIbYWQJY	236
FEWTSarGYWQJY	237
FEWTPGYWQPY	238
FEWTPGYWQHY	239
FEWTPGWYQJY	240

ACFEWTPGWYQJY 241 FEWTPGW-PY-QJY 242 FAWTPGYWQJY 243 FEWAPGYWQJY 244 FEWPGYWQJY 245 FEWTPGYWQJY 246 ACFEWTPGYWQJY 247 FEWTPAWYQJY 248 FEWTPSarWYQJY 249 FEWTPGYYQPY 250 FEWTPGWQPY 251 FEWTPGWWQPY 251 FEWTPGWWQPY 252 FEWTPRWQDY 253 FEWTPROGYWQJY 253 FEWTPAIDYWQJY 255 FEWTPAIDYWQJY 256 FEWTPGYWQJY 256 FEWTPGYWQJY 258 YEWTPGYYQJY 258 YEWTPGYYQJY 260 FEWTPSYYQJY 261 FEWTPNYYQJY 262 TKPR 263 RKSSK 264 RKQDK 265 NRKQDK 266 RKQDKR 266 NRKQDKRF 269 VTKFY 270
FAWTPGYWQJY 243 FEWAPGYWQJY 244 FEWYPGYWQJY 245 FEWTPGYWQJY 246 AcFEWTPGYWQJY 247 FEWTPAWYQJY 248 FEWTPSarWYQJY 249 FEWTPGYQPY 250 FEWTPGYWQPY 251 FEWTPGWQPY 251 FEWTPYWQJY 253 FEWTPYWQJY 253 FEWTPAIbYWQJY 254 FEWTPAIbYWQJY 255 FEWTSarGYWQJY 256 FEWTPGYWQPYALPL 257 INAPEWTPGYYQJY 258 YEWTPGYYQJY 260 FEWTPSYYQJY 260 FEWTPSYYQJY 261 FEWTPNYYQJY 262 TKPR 263 RKSK 264 RKQDK 265 NRKQDK 265 NRKQDK 265 NRKQDKRF 269 VTKFY 270 VTDFY 271 SHLYWQPYSVQ 673 <t< td=""></t<>
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FEWTPAWYQJY 248 FEWTPSarWYQJY 249 FEWTPGYYQPY 250 FEWTPGWWQPY 251 FEWTPGWWQPY 252 FEWTPNYWQJY 253 FEWTPecGYWQJY 254 FEWTPAIbYWQJY 255 FEWTSarGYWQJY 256 FEWTPGYWQPYALPL 257 1NapEWTPGYYQJY 258 YEWTPGYYQJY 260 FEWTPSYYQJY 261 FEWTPSYYQJY 262 TKPR 263 RKSK 264 RKQDK 265 NRKQDK 265 NRKQDK 266 RKQDKR 267 ENRKQDKRF 268 VTKFY 270 VTTKFY 271 SHLYWQPYSUQ 671 TLVYWQPYSUQT 672 RQDYWQPYSUQT 674 RLVYWQPYSUQT 675 SRVWFQPYSUQT 678 TFVYWQPYSUQR 676 TLVYWQPYSUQR 678
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DAYWVQ PYALPL	740
WSGYFQ PYALPL	741
NIEFWQ PYALPL	742
TROWVQ PYALPL	743
DSSWYQ PYALPL	744
IGNWYQ PYALPL	745
NLRWDQ PYALPL	746
LPEFWQ PYALPL	747
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RSQYYQ PYALPL	749
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NSYFWQ PYALPL	751
RFMYWQPYSVQR	752
AHLFWQPYSVQR	753
WWQPYALPL	754
YYQPYALPL	755
YFQPYALGL	756
YWYQPYALPL	757
RWWQPYATPL	758
GWYQPYALGF	759
YWYQPYALGL	760
IWYQPYAMPL	761
SNMQPYQRLS	762
TFVYWQPY AVGLPAAETACN	763
TFVYWQPY SVQMTITGKVTM	764
TFVYWQPY SSHXXVPXGFPL	765
TFVYWQPY YGNPQWAIHVRH	766
TFVYWQPY VLLELPEGAVRA	767
TFVYWQPY VDYVWPIPIAQV	768
GWYQPYVDGWR	769
RWEQPYVKDGWS	770
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LFEQPYAKALGL	773
GWEQPYARGLAG	774
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PWIQPYARGFG	779
RPLYWQPYSVQV	780
TLIYWQPYSVQI	781
RFDYWQPYSDQT	782
WHQFVQPYALPL	783
EWDS VYWQPYSVQ TLLR	784
WEQN VYWQPYSVQ SFAD	785
SDV VYWQPYSVQ SLEM	786
YYDG VYWQPYSVQ VMPA	787
SDIWYQ PYALPL	788
QRIWWQ PYALPL	789

Carrier Division But at Divisi	700
SRIWWQ PYALPL	790
RSLYWQ PYALPL	791 792
TIIWEQ PYALPL	
WETWYQ PYALPL	793 794
SYDWEQ PYALPL	
SRIWCQ PYALPL	795
EIMFWQ PYALPL	796
DYVWQQ PYALPL	797
MDLLVQ WYQPYALPL	798
GSKVIL WYQPYALPL	799
RQGANI WYQPYALPL	800
GGGDEP WYQPYALPL	801
SQLERT WYQPYALPL	802
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KKGSTQ WYQPYALPL	804
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VKQKWR WYQPYALPL	807
LRRHDV WYQPYALPL	808
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ESKEDQ WYQPYALPL	810
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EGSREG WYQPYALPL	812
VIEWWQ PYALPL	813
VWYWEQ PYALPL	814
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FYEWWQ PYALPL	816
EGWWVQ PYALPL	817
WGEWLQ PYALPL	818
DYVWEQ PYALPL	819
AHTWWQ PYALPL	820
FIEWFQ PYALPL	821
WLAWEQ PYALPL	822
VMEWWQ PYALPL	823
ERMWQ PYALPL	824
NXXWXX PYALPL	825
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TLYWEQ PYALPL	827
VWRWEQ PYALPL	828
LLWTQ PYALPL	829
SRIWXX PYALPL	830
SDIWYQ PYALPL	831
WGYYXX PYALPL	832
TSGWYQ PYALPL	833
VHPYXX PYALPL	834
EHSYFQ PYALPL	835
XXIWYQ PYALPL	836
AQLHSQ PYALPL	837
WANWFQ PYALPL	838
SRLYSQ PYALPL	839

SIVWSQ PYALPL 842 SRDLVQ PYALPL 842 HWGH VYWQPYSVQ DDLG 843 SWHS VYWQPYSVQ SVPE 844 WRDS VYWQPYSVQ SVPE 844 WRDS VYWQPYSVQ KWLD 846 TWDA VYWQPYSVQ SLDP 847 TWS VYWQPYSVQ SUPS 848 YWY QPY ALGL 849 YWY QPY ALPL 850 EWI QPY ATGL 851 NWE QPY AKPL 852 AFY QPY ALPL 853 FLY QPY ALPL 853 FLY QPY ALPL 854 VCK QPY LEWC 855 ETPFTWEESNAYYWQPYALPL 856 GGWLTWQDSVDMYWQPYALPL 857 FSEAGYTWPENTYWQPYALPL 858 TESPGGLDWAKIYWQPYALPL 860 TANVSSFEWTPGYWQPYALPL 861 SWGEDHNFWTSE YWQPYALPL 862 MNDQTSEVSTFP YWQPYALPL 862 MNDQTSEVSTFP YWQPYALPL 863 SWSEAFEQPRINL YWQPYALPL 865 MEKTYTTURDADL YWQPYALPL 866 THDEHI YWQPYALPL 867		
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SWSEAFEQPRNL YWQPYALPL QYAEPSALNDWG YWQPYALPL R65 NGDWATADWSNY YWQPYALPL R66 THDEHI YWQPYALPL R67 MLEKTYTTWTPG YWQPYALPL R68 WSDPLTRDADL YWQPYALPL SDAFTTQDSQAM YWQPYALPL R70 GDDAAWRTDSLT YWQPYALPL ENTYSPNWADSM YWQPYALPL S72 ENTYSPNWADSM YWQPYALPL SVGEDHNFWTSE YWQPYALPL SVGEDHNFWTSE YWQPYALPL R75 QTPFTWEESNAY YWQPYALPL R76 ENPFTWQESNAY YWQPYALPL R77 VTPFTWEDSNVF YWQPYALPL R78 QAPLTWQESAAY YWQPYALPL R89 EPTFTWEESKAT YWQPYALPL R80 EPTFTWEESNAY YWQPYALPL R81 TTLLTWEESNAY YWQPYALPL R82 ESPLTWEESNAY YWQPYALPL R83 ETPLTWEESNAY YWQPYALPL R84 EAFTWAESNAY YWQPYALPL R85 EALFTWKESTAY YWQPYALPL R86 STP-TWEESNAY YWQPYALPL R87 EALFTWKESTAY YWQPYALPL R86 STP-TWEESNAY YWQPYALPL R87 ETPFTWEESNAY YWQPYALPL R87		862
QYAEPSALNDWG YWQPYALPL NGDWATADWSNY YWQPYALPL 866 THDEHI YWQPYALPL 867 MLEKTYTTWTPG YWQPYALPL 868 WSDPLTRDADL YWQPYALPL 869 SDAFTTQDSQAM YWQPYALPL 870 GDDAAWRTDSLT YWQPYALPL 871 AIIRQLYRWSEM YWQPYALPL ENTYSPNWADSM YWQPYALPL 872 ENTYSPNWADSM YWQPYALPL 873 MNDQTSEVSTFP YWQPYALPL 874 SVGEDHNFWTSE YWQPYALPL 875 QTPFTWEESNAY YWQPYALPL 877 VTPFTWEDSNVF YWQPYALPL 878 QIPFTWQSNAY YWQPYALPL 879 QAPLTWQESAAY YWQPYALPL 880 EPTFTWEESKAT YWQPYALPL 881 TTTLTWEESNAY YWQPYALPL 882 ESPLTWEESSAL YWQPYALPL 883 ETPLTWEESNAY YWQPYALPL 884 EAFTWAESNAY YWQPYALPL 885 EALFTWKESTAY YWQPYALPL 886 STP-TWEESNAY YWQPYALPL 887 ETPFTWEESNAY YWQPYALPL 887		863
NGDWATADWSNY YWQPYALPL THDEHI YWQPYALPL MLEKTYTTWTPG YWQPYALPL WSDPLTRDADL YWQPYALPL SDAFTTQDSQAM YWQPYALPL B70 GDDAAWRTDSLT YWQPYALPL AIIRQLYRWSEM YWQPYALPL ENTYSPNWADSM YWQPYALPL S72 ENTYSPNWADSM YWQPYALPL S73 MNDQTSEVSTFP YWQPYALPL S75 QTPFTWEESNAY YWQPYALPL ENPFTWQESNAY YWQPYALPL B76 ENPFTWQESNAY YWQPYALPL B77 VTPFTWEDSNVF YWQPYALPL B78 QIPFTWEGSNAY YWQPYALPL B79 QAPLTWQESAAY YWQPYALPL B80 EPTFTWEESKAT YWQPYALPL B81 TTTLTWEESNAY YWQPYALPL B82 ESPLTWEESSAL YWQPYALPL B83 ETPLTWEESNAY YWQPYALPL B84 EATFTWAESNAY YWQPYALPL B85 EALFTWKESTAY YWQPYALPL B86 STP-TWEESNAY YWQPYALPL B87 ETPLTWEESNAY YWQPYALPL B86 STP-TWEESNAY YWQPYALPL B87 ETPLTWEESNAY YWQPYALPL B86 STP-TWEESNAY YWQPYALPL B87 ETPFTWEESNAY YWQPYALPL B87		864
THDEHI YWQPYALPL MLEKTYTTWTPG YWQPYALPL 868 WSDPLTRDADL YWQPYALPL SDAFTTQDSQAM YWQPYALPL GDDAAWRTDSLT YWQPYALPL AIIRQLYRWSEM YWQPYALPL ENTYSPNWADSM YWQPYALPL 872 ENTYSPNWADSM YWQPYALPL 873 MNDQTSEVSTFP YWQPYALPL 874 SVGEDHNFWTSE YWQPYALPL 875 QTPFTWEESNAY YWQPYALPL 877 VTPFTWEESNAY YWQPYALPL 877 QAPLTWQESAAY YWQPYALPL 880 EPTFTWEESKAT YWQPYALPL 881 TTTLTWEESNAY YWQPYALPL 882 ESPLTWEESSAL YWQPYALPL 883 ETPLTWEESNAY YWQPYALPL 884 EATFTWAESNAY YWQPYALPL 885 EALFTWKESTAY YWQPYALPL 886 STP-TWEESNAY YWQPYALPL 887 ETPLTWEESNAY YWQPYALPL 887 ETPLTWEESNAY YWQPYALPL 888		
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SDAFTTQDSQAM YWQPYALPL GDDAAWRTDSLT YWQPYALPL AIIRQLYRWSEM YWQPYALPL ENTYSPNWADSM YWQPYALPL ENTYSPNWADSM YWQPYALPL 873 MNDQTSEVSTFP YWQPYALPL 874 SVGEDHNFWTSE YWQPYALPL 875 QTPFTWEESNAY YWQPYALPL 877 VTPFTWEDSNVF YWQPYALPL 877 VTPFTWEDSNVF YWQPYALPL 879 QAPLTWQESAAY YWQPYALPL 880 EPTFTWEESKAT YWQPYALPL 881 TTTLTWEESNAY YWQPYALPL 882 ESPLTWEESNAY YWQPYALPL 883 ETPLTWEESNAY YWQPYALPL 884 EATFTWAESNAY YWQPYALPL 885 EALFTWKESTAY YWQPYALPL 886 STP-TWEESNAY YWQPYALPL 887 ETPFTWEESNAY YWQPYALPL 887		868
GDDAAWRTDSLT YWQPYALPL AIIRQLYRWSEM YWQPYALPL ENTYSPNWADSM YWQPYALPL 873 MNDQTSEVSTFP YWQPYALPL 874 SVGEDHNFWTSE YWQPYALPL 875 QTPFTWEESNAY YWQPYALPL 876 ENPFTWQESNAY YWQPYALPL 877 VTPFTWEDSNVF YWQPYALPL 879 QAPLTWQESAAY YWQPYALPL 880 EPTFTWEESKAT YWQPYALPL 881 TTTLTWEESNAY YWQPYALPL 882 ESPLTWEESSAL YWQPYALPL 883 ETPLTWEESNAY YWQPYALPL 884 EATFTWAESNAY YWQPYALPL 885 EALFTWKESTAY YWQPYALPL 886 STP-TWEESNAY YWQPYALPL 887 ETPFTWEESNAY YWQPYALPL 887		
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MNDQTSEVSTFP YWQPYALPL SVGEDHNFWTSE YWQPYALPL R76 QTPFTWEESNAY YWQPYALPL ENPFTWQESNAY YWQPYALPL 877 VTPFTWEDSNVF YWQPYALPL R79 QIPFTWEQSNAY YWQPYALPL 880 EPTFTWEESKAT YWQPYALPL 881 TTTLTWEESNAY YWQPYALPL 882 ESPLTWEESSAL YWQPYALPL 883 ETPLTWEESNAY YWQPYALPL 884 EATFTWAESNAY YWQPYALPL 885 EALFTWKESTAY YWQPYALPL 886 STP-TWEESNAY YWQPYALPL 887 ETPFTWEESNAY YWQPYALPL 887	AIIRQLYRWSEM YWQPYALPL	872
SVGEDHNFWTSE YWQPYALPL QTPFTWEESNAY YWQPYALPL ENPFTWQESNAY YWQPYALPL 877 VTPFTWEDSNVF YWQPYALPL QIPFTWEQSNAY YWQPYALPL 879 QAPLTWQESAAY YWQPYALPL 880 EPTFTWEESKAT YWQPYALPL 881 TTTLTWEESNAY YWQPYALPL 882 ESPLTWEESSAL YWQPYALPL 883 ETPLTWEESNAY YWQPYALPL 884 EATFTWAESNAY YWQPYALPL 885 EALFTWKESTAY YWQPYALPL 886 STP-TWEESNAY YWQPYALPL 887 ETPFTWEESNAY YWQPYALPL 887	ENTYSPNWADSM YWQPYALPL	873
QTPFTWEESNAY YWQPYALPL 876 ENPFTWQESNAY YWQPYALPL 877 VTPFTWEDSNVF YWQPYALPL 878 QIPFTWEQSNAY YWQPYALPL 879 QAPLTWQESAAY YWQPYALPL 880 EPTFTWEESKAT YWQPYALPL 881 TTTLTWEESNAY YWQPYALPL 882 ESPLTWEESSAL YWQPYALPL 883 ETPLTWEESNAY YWQPYALPL 884 EATFTWAESNAY YWQPYALPL 885 EALFTWKESTAY YWQPYALPL 885 EALFTWKESTAY YWQPYALPL 886 STP-TWEESNAY YWQPYALPL 887 ETPFTWEESNAY YWQPYALPL 887	MNDQTSEVSTFP YWQPYALPL	874
ENPFTWQESNAY YWQPYALPL 877 VTPFTWEDSNVF YWQPYALPL 878 QIPFTWEQSNAY YWQPYALPL 879 QAPLTWQESAAY YWQPYALPL 880 EPTFTWEESKAT YWQPYALPL 881 TTTLTWEESNAY YWQPYALPL 882 ESPLTWEESSAL YWQPYALPL 883 ETPLTWEESNAY YWQPYALPL 884 EATFTWAESNAY YWQPYALPL 885 EALFTWKESTAY YWQPYALPL 886 STP-TWEESNAY YWQPYALPL 887 ETPFTWEESNAY YWQPYALPL 888	SVGEDHNFWTSE YWQPYALPL	875
VTPFTWEDSNVF YWQPYALPL 878 QIPFTWEQSNAY YWQPYALPL 879 QAPLTWQESAAY YWQPYALPL 880 EPTFTWEESKAT YWQPYALPL 881 TTTLTWEESNAY YWQPYALPL 882 ESPLTWEESSAL YWQPYALPL 883 ETPLTWEESNAY YWQPYALPL 884 EATFTWAESNAY YWQPYALPL 885 EALFTWKESTAY YWQPYALPL 886 STP-TWEESNAY YWQPYALPL 886 STP-TWEESNAY YWQPYALPL 887 ETPFTWEESNAY YWQPYALPL 887	QTPFTWEESNAY YWQPYALPL	
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ETPFTWEESNAY YWQPYALPL 888	EALFTWKESTAY YWQPYALPL	886
	STP-TWEESNAY YWQPYALPL	887
VADETNIESCAV VINORVALDI 880	ETPFTWEESNAY YWQPYALPL	888
RAFFIWEESGAT TWGFTALFE 009	KAPFTWEESQAY YWQPYALPL	889

LOZOFTHIE CONAVAMORYAL DI	200
STSFTWEESNAY YWQPYALPL	890 891
DSTFTWEESNAY YWQPYALPL	892
YIPFTWEESNAY YWQPYALPL	
QTAFTWEESNAY YWQPYALPL	893
ETLFTWEESNAT YWQPYALPL	894
VSSFTWEESNAY YWQPYALPL	895
QPYALPL	896
Py-1-NapPYQJYALPL	897
TANVSSFEWTPG YWQPYALPL	898
FEWTPGYWQPYALPL	899
FEWTPGYWQJYALPL	900
FEWTPGYYQJYALPL	901
ETPFTWEESNAYYWQPYALPL	902
FTWEESNAYYWQJYALPL	903
ADVL YWQPYA PVTLWV	904
GDVAE YWQPYA LPLTSL	905
SWTDYG YWQPYA LPISGL	906
FEWTPGYWQPYALPL	911
FEWTPGYWQJYALPL	912
FEWTPGWYQPYALPL	913
FEWTPGWYQJYALPL	914
FEWTPGYYQPYALPL	915
FEWTPGYYQJYALPL	916
TANVSSFEWTPGYWQPYALPL	918
SWTDYGYWQPYALPISGL	919
ETPFTWEESNAYYWQPYALPL	920
ENTYSPNWADSMYWQPYALPL	921
SVGEDHNFWTSEYWQPYALPL	922
DGYDRWRQSGERYWQPYALPL	923
FEWTPGYWQPYALPL	924
FEWTPGYWQPY	925
FEWTPGYWQJY	926
EWTPGYWQPY	927
FEWTPGWYQJY	928
AEWTPGYWQJY	929
FAWTPGYWQJY	930
FEATPGYWQJY	931
FEWAPGYWQJY	932
FEWTAGYWQJY	933
FEWTPAYWQJY	934
FEWTPGAWQJY	935
FEWTPGYAQJY	936
FEWTPGYWQJA	937
FEWTGGYWQJY	938
FEWTPGYWQJY	939
FEWTJGYWQJY	940
FEWTPecGYWQJY	941
FEWTPAIDYWQJY	942
FEWTPSarWYQJY	943
FEWTSarGYWQJY	944
[

FEWTPNYWQJY	945
FEWTPVYWQJY	946
FEWTVPYWQJY	947
AcFEWTPGWYQJY	948
AcFEWTPGYWQJY	949
INap-EWTPGYYQJY	950
YEWTPGYYQJY	951
FEWVPGYYQJY	952
FEWTPGYYQJY	953
FEWTPSYYQJY	954
FEWTPnYYQJY	955
SHLY-Nap-QPYSVQM	956
TLVY-Nap-QPYSLQT	957
RGDY-Nap-QPYSVQS	958
NMVY-Nap-QPYSIQT	959
VYWQPYSVQ	960
VY-Nap-QPYSVQ	961
TFVYWQJYALPL	962
FEWTPGYYQJ-Bpa	963
XaaFEWTPGYYQJ-Bpa	964
FEWTPGY-Bpa-QJY	965
AcFEWTPGY-Bpa-QJY	966
FEWTPG-Bpa-YQJY	967
AcFEWTPG-Bpa-YQJY	968
AcFE-Bpa-TPGYYQJY	969
AcFE-Bpa-TPGYYQJY	970
Bpa-EWTPGYYQJY	971
AcBpa-EWTPGYYQJY	972
VYWQPYSVQ	973
RLVYWQPYSVQR	974
RLVY-Nap-QPYSVQR	9 7 5
RLDYWQPYSVQR	976
RLVWFQPYSVQR	977
RLVYWQPYSIQR	978
DNSSWYDSFLL	980
DNTAWYESFLA	981
DNTAWYENFLL	982
PARE DNTAWYDSFLI WC	983
TSEY DNTTWYEKFLA SQ	984
SQIP DNTAWYQSFLL HG	985
SPFI DNTAWYENFLL TY	986
EQIY DNTAWYDHFLL SY	987
TPFI DNTAWYENFLL TY	988
TYTY DNTAWYERFLM SY	989
TMTQ DNTAWYENFLL SY	990
TI DNTAWYANLVQ TYPQ	991
TI DNTAWYERFLA QYPD	992
HI DNTAWYENFLL TYTP	993
SQ DNTAWYENFLL SYKA	994
QI DNTAWYERFLL QYNA	995
2/-	

NQ DNTAWYESFLL QYNT	996
TI DNTAWYENFLL NHNL	997
HY DNTAWYERFLQ QGWH	998
ETPFTWEESNAYYWQPYALPL	999
YIPFTWEESNAYYWQPYALPL	1000
DGYDRWRQSGERYWQPYALPL	1001
pY-INap-pY-QJYALPL	1002
TANVSSFEWTPGYWQPYALPL	1003
FEWTPGYWQJYALPL	1004
FEWTPGYWQPYALPLSD	1005
FEWTPGYYQJYALPL	1006
FEWTPGYWQJY	1007
AcFEWTPGYWQJY	1008
AcFEWTPGWYQJY	1009
AcFEWTPGYYQJY	1010
AcFEWTPaYWQJY	1011
AcFEWTPaWYQJY	1012
AcFEWTPaYYQJY	1013
FEWTPGYYQJYALPL	1014
FEWTPGYWQJYALPL	1015
FEWTPGWYQJYALPL	1016
TANVSSFEWTPGYWQPYALPL	1017
AcFEWTPGYWQJY	1018
AcFEWTPGWYQJY	1019
AcFEWTPGYYQJY	1020
AcFEWTPAYWQJY	1021
ACFEWTPAWYQJY	1022
AcFEWTPAYYQJY	1023

Table 5—EPO-mimetic peptide sequences

	CEO
Sequence/structure	SEQ
NA CANADA CENTA CANADA	ID NO:
YXCXXGPXTWXCXP	63
YXCXXGPXTWXCXP-YXCXXGPXTWXCXP	84
YXCXXGPXTWXCXP-A-YXCXXGPXTWXCXP	85
YXCXXGPXTWXCXP-Λ- (ε-amine)	86
βA	0,
YXCXXGPXTWXCXP-Λ- (α-amine)	86
GGTYSCHFGPLTWVCKPQGG	87
GGDYHCRMGPLTWVCKPLGG	88
GGVYACRMGPITWVCSPLGG	89
VGNYMCHFGPITWVCRPGGG	90
GGLYLCRFGPVTWDCGYKGG	91
GGTYSCHFGPLTWVCKPQGG- GGTYSCHFGPLTWVCKPQGG	92
GGTYSCHFGPLTWVCKPQGG -A-	93
GGTYSCHFGPLTWVCKPQGG	
GGTYSCHFGPLTWVCKPQGGSSK	94
GGTYSCHFGPLTWVCKPQGGSSK- GGTYSCHFGPLTWVCKPQGGSSK	95
GGTYSCHFGPLTWVCKPQGGSSK-A-	96
GGTYSCHFGPLTWVCKPQGGSSK GGTYSCHFGPLTWVCKPQGGSS	97
(ε-amine)	
βA GGTYSCHFGPLTWVCKPQGGSS (α-amine)	97
GGTYSCHFGPLTWVCKPQGGSSK(-A-biotin)	98
CX,X,GPX,TWX,C	421
GGTYSCHGPLTWVCKPQGG	422
VGNYMAHMGPITWVCRPGG	423
GGPHHVYACRMGPLTWIC	424
GGTYSCHFGPLTWVCKPQ	425
GGLYACHMGPMTWVCQPLRG	_ 426
TIAQYICYMGPETWECRPSPKA	427
YSCHFGPLTWVCK	428
YCHFGPLTWVC	429
X ₃ X ₄ X ₅ GPX ₆ TWX ₇ X ₈	124
YX ₂ X ₃ X ₄ X ₅ GPX ₅ TWX ₇ X ₈	461

X,YX,X,X,GPX,TWX,X,X,X,X,1	419
X,YX,CX,X,GPX,TWX,CX,X,,X,,	420
GGLYLCRFGPVTWDCGYKGG	1024
GGTYSCHFGPLTWVCKPQGG	1025
GGDYHCRMGPLTWVCKPLGG	1026
VGNYMCHFGPITWVCRPGGG	1029
GGVYACRMGPITWVCSPLGG	1030
VGNYMAHMGPITWVCRPGG	1035
GGTYSCHFGPLTWVCKPQ	1036
GGLYACHMGPMTWVCQPLRG	1037
TIAQYICYMGPETWECRPSPKA	1038
YSCHFGPLTWVCK	1039
YCHFGPLTWVC	1040
SCHFGPLTWVCK	1041
(AX ₂) _n X ₃ X ₄ X ₅ GPX ₅ TWX ₇ X ₈	1042

Table 6—TPO-mimetic peptide sequences

Sequence/structure	SEQ ID NO:
IEGPTLRQWLAARA	13
IEGPTLRQWLAAKA	24
IEGPTLREWLAARA	25
IEGPTLRQWLAARA-A-IEGPTLRQWLAARA	26
IEGPTLRQWLAAKA-A-IEGPTLRQWLAAKA	27
IEGPTLRQCLAARA-A-IEGPTLRQCLAARA	28
IEGPTLRQWLAARA-A-K(BrAc)-A-IEGPTLRQWLAARA	29
IEGPTLRQWLAARA-A-K(PEG)-A-IEGPTLRQWLAARA	30
IEGPTLRQCLAARA-A-IEGPTLRQWLAARA	31
IEGPTLRQCLAARA-A-IEGPTLRQWLAARA	31
IEGPTLRQWLAARA-A-IEGPTLRQCLAARA	32
IEGPTLRQWLAARA-A-IEGPTLRQCLAARA	32
VRDQIXXXL	33
TLREWL	34
GRVRDQVAGW	35
GRVKDQIAQL	36
GVRDQVSWAL	37
ESVREQVMKY	38
SVRSQISASL	39
GVRETVYRHM	40
GVREVIVMHML	41
GRVRDQIWAAL	42
AGVRDQILIWL	43
GRVRDQIMLSL	44
GRVRDQI(X) ₃ L	45
CTLRQWLQGC	46
CTLQEFLEGC	47
CTRTEWLHGC	48
CTLREWLHGGFC	49
CTLREWVFAGLC	50
CTLRQWLILLGMC	51
CTLAEFLASGVEQC	52
CSLQEFLSHGGYVC	53
CTLREFLDPTTAVC	54
CTLKEWLVSHEVWC	55
CTLREWL(X) ₂₈ C	56-60
REGPTLRQWM	61
EGPTLRQWLA	62
ERGPFWAKAC	63
REGPRCVMWM	64
CGTEGPTLSTWLDC	65

CEQDGPTLLEWLKC	66
CELVGPSLMSWLTC	67
CLTGPFVTQWLYEC	68
CRAGPTLLEWLTLC	69
CADGPTLREWISFC	70
C(X), EGPTLREWL(X), C	71-74
GGCTLREWLHGGFCGG	75
GGCADGPTLREWISFCGG	76
GNADGPTLRQWLEGRRPKN	77
LAIEGPTLRQWLHGNGRDT	78
HGRVGPTLREWKTQVATKK	79
TIKGPTLRQWLKSREHTS	80
ISDGPTLKEWLSVTRGAS	81
SIEGPTLREWLTSRTPHS	82

Table 7—G-CSF-mimetic peptide sequences

Sequence/structure	SEQ
_	ID NO:
EEDCK	99
EEDCK	99
EEDCK	99
EEDøK	100
EEDøK	100
1	<u>.</u> [
EEDoK	100
pGluEDσK	101
pGluEDσK	101
pGluEDoK	101
PicSDσK	102
PicSDoK	102
PicSDoK	102
EEDCK-A-EEDCK	103
EEDXK-A-EEDXK	104

Table 8—TNF-antagonist peptide sequences

Sequence/structure	SEQ
1	ID NO:
YCFTASENHCY	106
YCFTNSENHCY	107
YCFTRSENHCY	108
FCASENHCY	109
YCASENHCY	110
FCNSENHCY	111
FCNSENRCY	112
FCNSVENRCY	113
YCSQSVSNDCF	114
FCVSNDRCY	115
YCRKELGQVCY	116
YCKEPGQCY	117
YCRKEMGCY	118
FCRKEMGCY	119
YCWSQNLCY	120
YCELSQYLCY	121
YCWSQNYCY	122
YCWSQYLCY	123
DFLPHYKNTSLGHRP	1085
AA,-AB,	NR
` \	
AC	
/	
AA,-AB,	

Table 9—Integrin-binding peptide sequences

Sequence/structure	SEQ
Sequence/structure	ID NO:
RX,ETX,WX,	441
RX,ETX,WX,	442
RGDGX	443
CRGDGXC	444
	445
CX,X,RLDX,X,C CARRLDAPC	446
CPSRLDSPC	447
	448
X,X,X,RGDX,X,X, CX,CRGDCX,C	449
CDCRGDCFC	450
CDCRGDCLC	451
CLCRGDCIC	452
	453
X,X,DDX,X,X,X	454
X,X,X,DDX,X,X,X,X, CWDDGWLC	455
CWDDLWWLC	456
CWDDGLMC	457
	458
CWDDGWMC	459
CSWDDGWLC CPDDLWWLC	460
	NR NR
NGR	NR NR
GSL	NR NR
RGD CGRECPRLCQSSC	1071
	1072
CNGRCVSGCAGRC	1073
CLSGSLSC RGD	NR
	NR NR
NGR GSL	NR NR
	1074
NGRAHA	1075
CDCRGDCFC	1076
CGSLVRC	1070
DLXXL	1043
	1043
RTDLDSLRTYTL RTDLDSLRTY	1053
RTDLDSLRT	1054
RTDLDSLR	1078
GDLDLLKLRLTL	1079
GDLHSLRQLLSR	1079
RDDLHMLRLQLW	1081
SSDLHALKKRYG	1082
RGDLKQLSELTW	1083
RGDLAALSAPPV	1084
RGDLAALSAFFV	1004

Table 10—Selectin antagonist peptide sequences

Sequence/structure	SEQ
-	ID NO:
DITWDQLWDLMK	147
DITWDELWKIMN	148
DYTWFELWDMMQ	149
QITWAQLWNMMK	150
DMTWHDLWTLMS	151
DYSWHDLWEMMS	152
EITWDQLWEVMN	153
HVSWEQLWDIMN	154
HITWDQLWRIMT	155
RNMSWLELWEHMK	156
AEWTWDQLWHVMNPAESQ	157
HRAEWLALWEQMSP	158
KKEDWLALWRIMSV	159
ITWDQLWDLMK	160
DITWDQLWDLMK	161
DITWDQLWDLMK	162
DITWDQLWDLMK	163
CQNRYTDLVAIQNKNE	462
AENWADNEPNNKRNNED	463
RKNNKTWTWVGTKKALTNE	464
KKALTNEAENWAD	465
CQXRYTDLVAIQNKXE	466
RKXNXXWTWVGTXKXLTEE	467
AENWADGEPNNKXNXED	468
CXXXYTXLVAIQNKXE	469
RKXXXXWXWVGTXKXLTXE	470
AXNWXXXEPNNXXXED	471
XKXKTXEAXNWXX	472

Table 11—Antipathogenic peptide sequences

C	SEQ
Sequence/structure	ID NO:
OFFALIBRIESERI FIXTUL SAVOSAL SSSGGOO	503
GFFALIPKIISSPLFKTLLSAVGSALSSSGGQQ	504
GFFALIPKIISSPLFKTLLSAVGSALSSSGGQE	505
GFFALIPKIISSPLFKTLLSAV	506
GFFALIPKIISSPLFKTLLSAV	507
KGFFALIPKIISSPLFKTLLSAV	508
KKGFFALIPKIISSPLFKTLLSAV KKGFFALIPKIISSPLFKTLLSAV	509
	510
GFFALIPKIIS GIGAVLKVLTTGLPALISWIKRKRQQ	511
	512
GIGAVLKVLTTGLPALISWIKRKRQQ	513
GIGAVLKVLTTGLPALISWIKRKRQQ	
GIGAVLKVLTTGLPALISWIKR	514 515
AVLKVLTTGLPALISWIKR	
KLLLLIKLLLK	516
KLLLKLLK	517
KLLLKLKLLK	518
KKLLKLKLKK	519
KLLLKLLKLLK	520
KLLLKLKLKLK	521
KLLLLK	522
KLLLKLLK	523
KLLLKLKLKLK	524
KLLLKLKLKLK	525
KLLKLKLKLK	526
KAAAKAAKAAK	527
KVVVKVVKVVK	528
KVVVKVKVKVVK	529
KVVVKVKVKVK	530
KVVVKVKVKVVK	531
KLILKL	532
KVLHLL	533
LKLRLL	534
KPLHLL	535
KLILKLVR	536
KVFHLLHL	537
HKFRILKL	538
KPFHILHL	539
KIIIKIKIKIK	540
KIIIKIKIKI	541
KIIIKIKIKIK	542
KIPIKIKIKIPK	543
KIPIKIKIVK	544
RIIIRIRIRI	545
RIIIRIRIRIR	546
RIIIRIRIRIR	547
RIVIRIRIRLIR	548

	T 740
RIIVRIRLRIIR	549
RIGIRLRVRIIR	550
KIVIRIRIRLIR	551
RIAVKWRLRFIK	552
KIGWKLRVRIIR	553
KKIGWLIIRVRR	554
RIVIRIRIRIR	555
RIIVRIRLRIIRVR	556
RIGIRLRVRIIRRV	557
KIVIRIRARLIRIRIR	558
RIIVKIRLRIIKKIRL	559
KIGIKARVRIIRVKII	560
RIIVHIRLRIIHHIRL	561
HIGIKAHVRIIRVHII	562
RIYVKIHLRYIKKIRL	563
KIGHKARVHIIRYKII	564
RIYVKPHPRYIKKIRL	565
KPGHKARPHIIRYKII	566
KIVIRIRIRIRIRKIV	567
RIIVKIRLRIIKKIRLIKK	568
KIGWKLRVRIIRVKIGRLR	569
KIVIRIRIRIRIRIKIVKVKRIR	570
RFAVKIRLRIIKKIRLIKKIRKRVIK	571
KAGWKLRVRIIRVKIGRLRKIGWKKRVRIK	572
RIYVKPHPRYIKKIRL	573
KPGHKARPHIIRYKII	574
KIVIRIRIRIRIRKIV	575
RIIVKIRLRIIKKIRLIKK	576
RIYVSKISIYIKKIRL	577
KIVIFTRIRLTSIRIRSIV	578
KPIHKARPTIIRYKMI	579
cyclicCKGFFALIPKIISSPLFKTLLSAVC	580
CKKGFFALIPKIISSPLFKTLLSAVC	581
CKKKGFFALIPKIISSPLFKTLLSAVC	582
CyclicCRIVIRIRIRLIRIRC	583
CyclicCKPGHKARPHIIRYKIIC	584
CyclicCRFAVKIRLRIIKKIRLIKKIRKRVIKC	585
KLLLKLLL KLLKC	586
KLLLKLLKLLK	587
KLLLKLKLKC	588
KLLLKLLK	589

Table 12—VIP-mimetic peptide sequences

Sequence/structure	SEQ
•	ID NO:
HSDAVFYDNYTR LRKQMAVKKYLN SILN	590
NIE HSDAVFYDNYTR LRKQMAVKKYLN SILN	591
X, X, 'X, " X,	592
X, S X, LN	593
NH CH CO KKYX5 NH CH CO X6	594
(CH2)mZ(CH2)n	
KKYL	595
NSILN	596
KKYL	597
KKYA	598
AVKKYL	599
NSILN	600
KKYV	601
SILauN	602
KKYLNIe	603
NSYLN	604
NSIYN	605
KKYLPPNSILN	606
LauKKYL	607
СарККҮL	608
KYL	NR
KKYNie	609
VKKYL	610
LNSILN	611
YLNSILN	612
KKYLN	613
KKYLNS	614
KKYLNSI	615
KKYLNSIL	616
KKYL	617
KKYDA	618
AVKKYL	619
NSILN	620
KKYV	621
SILauN	622
NSYLN	623
NSIYN	625
KKYLNIe	625
KKYLPPNSILN	626
KKYL	
KKYDA	628 -
AVKKYL	630
NSILN	631
KKYV	632
SILauN	1 034

[633
LauKKYL	634
CapKKYL	NR
KYL	NR NR
KYL	635
KKYNle	636
VKKYL	
LNSILN	637
YLNSILN	638
KKYLNle	639
KKYLN	640
KKYLNS	641
KKYLNSI	642
KKYLNSIL	643
KKKYLD	644
cyclicCKKYLC	645
CKKYLK	646
]	
S-CH ₂ -CO	
KKYA	647
WWTDTGLW	648
WWTDDGLW	649
WWDTRGLWVWTI	650
FWGNDGIWLESG	651
DWDQFGLWRGAA	652
RWDDNGLWVVVL	653
SGMWSHYGIWMG	654
GGRWDQAGLWVA	655
KLWSEQGIWMGE	656
CWSMHGLWLC	657
GCWDNTGIWVPC	658
DWDTRGLWVY	659
SLWDENGAWI	660
KWDDRGLWMH	661
QAWNERGLWT	662
QWDTRGLWVA	663
WNVHGIWQE	664
SWDTRGLWVE	665
DWDTRGLWVA	666
SWGRDGLWIE	667
EWTDNGLWAL	668
SWDEKGLWSA	669
SWDSSGLWMD	670

Table 13—Mdm/hdm antagonist peptide sequences

Sequence/structure	SEQ
•	ID NO:
TFSDLW	130
QETFSDLWKLLP	131
QPTFSDLWKLLP	· 132
QETFSDYWKLLP	133
QPTFSDYWKLLP	134
MPRFMDYWEGLN	135
VQNFIDYWTQQF	136
TGPAFTHYWATF	137
IDRAPTFRDHWFALV	138
PRPALVFADYWETLY	139
PAFSRFWSDLSAGAH	140
PAFSRFWSKLSAGAH	141
PXFXDYWXXL	142
QETFSDLWKLLP	143
QPTFSDLWKLLP	144
QETFSDYWKLLP	145
QPTFSDYWKLLP	146

Table 14—Calmodulin antagonist peptide sequences

Sequence/structure	SEQ ID NO:
SCVKWGKKEFCGS	164
SCWKYWGKECGS	165
SCYEWGKLRWCGS	166
SCLRWGKWSNCGS	167
SCWRWGKYQICGS	168
SCVSWGALKLCGS	169
SCIRWGQNTFCGS	170
SCWQWGNLKICGS	171
SCVRWGQLSICGS	172
LKKFNARRKLKGAILTTMLAK	173
RRWKKNFIAVSAANRFKK	174
RKWQKTGHAVRAIGRLSS	175
INLKALAALAKKIL	176
KIWSILAPLGTTLVKLVA	177
LKKLLKLLKKL	178
LKWKKLLKLLKKLL	179
AEWPSLTEIKTLSHFSV	180
AEWPSPTRVISTTYFGS	181
AELAHWPPVKTVLRSFT	182 -
AEGSWLQLLNLMKQMNN	183
AEWPSLTEIK	184

Table 15—Mast cell antagonists/Mast cell protease inhibitor peptide sequences

Sequence/structure	SEQ
	ID NO:
SGSGVLKRPLPILPVTR	272
RWLSSRPLPPLPLPPRT	273
GSGSYDTLALPSLPLHPMSS	274
GSGSYDTRALPSLPLHPMSS	275
GSGSSGVTMYPKLPPHWSMA	276
GSGSSGVRMYPKLPPHWSMA	277
GSGSSSMRMVPTIPGSAKHG	278
RNR	NR
QT	NR
RQK	NR
NRQ	NR
RQK	NR
RNRQKT	436
RNRQ	437
RNRQK	438
NRQKT	439
RQKT	440

Table 16—SH3 antagonist peptide sequences

Sequence/structure	SEQ
•	ID NO:
RPLPPLP	282
RELPPLP	283
SPLPPLP	284
GPLPPLP	· 285
RPLPIPP	286
RPLPIPP	287
RRLPPTP	288
RQLPPTP	289
RPLPSRP	290
RPLPTRP	291
SRLPPLP	292
RALPSPP	293
RRLPRTP	294
RPVPPIT	295
ILAPPVP	296
RPLPMLP	297
RPLPILP	298
RPLPSLP	299
RPLPSLP	300
RPLPMIP	301
RPLPLIP	302
RPLPPTP	303
RSLPPLP	304
RPQPPPP	305
RQLPIPP	306
XXXRPLPPLPXP	307
XXXRPLPPIPXX	308
XXXRPLPPLPXX	309
RXXRPLPPLPXP	310
RXXRPLPPLPPP	311
PPPYPPPIPXX	312
PPPYPPPVPXX	313
LXXRPLPXYP	314
ΨXXRPLPXLP	315
РРХӨХРРРҰР	316
+PPYPXKPXWL	317
RPXYPYR+SXP	318
PPVPPRPXXTL	319
ЧР Ч Г РЧК	320
+@DXPLPXLP	321

Table 17—Somatostatin or cortistatin mimetic peptide sequences

Sequence/structure	SEQ ID NO:
X¹-X²-Asn-Phe-Phe-Trp-Lys-Thr-Phe-X³-Ser-X⁴	473
Asp Arg Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys Lys	474
Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys Lys	475
Cys Arg Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys Lys	476
Asp Arg Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys	477
Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys	478
Cys Arg Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys	479
Asp Arg Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys	480
Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys Lys	481
Cys Lys Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys Lys	482
Asp Arg Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys	483
Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys	484
Cys Lys Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys	485
Asp Arg Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys Lys	486
Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys Lys	487
Cys Arg Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys Lys	488
Asp Arg Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys	489
Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys	490
Cys Arg Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys	491
Asp Arg Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys Lys	492
Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys Lys	493
Cys Lys Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys Lys	494
Asp Arg Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys	495
Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys	496
Cys Lys Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys	497

Table 18—UKR antagonist peptide sequences

Sequence/structure	SEQ ID NO:
AEPMPHSLNFSQYLWYT	196
AEHTYSSLWDTYSPLAF	197
AELDLWMRHYPLSFSNR	198
AESSLWTRYAWPSMPSY	199
AEWHPGLSFGSYLWSKT	200
AEPALLNWSFFFNPGLH	201
AEWSFYNLHLPEPQTIF	202
AEPLDLWSLYSLPPLAM	203
AEPTLWQLYQFPLRLSG	204
AEISFSELMWLRSTPAF	205
AELSEADLWTTWFGMGS	206
AESSLWRIFSPSALMMS	207
AESLPTLTSILWGKESV	208
AETLFMDLWHDKHILLT	209
AEILNFPLWHEPLWSTE	210
AESQTGTLNTLFWNTLR	211
AEPVYQYELDSYLRSYY	430
AELDLSTFYDIQYLLRT	431
AEFFKLGPNGYVYLHSA	432
FKLXXXGYVYL	433
AESTYHHLSLGYMYTLN	434
YHXLXXGYMYT	435

Table 19—Macrophage and/or
T-cell inhibiting peptide sequences

Sequence/structure	SEQ
•	ID NO:
Xaa-Yaa-Arg	NR
Arg-Yaa-Xaa	NR
Xaa-Arg-Yaa	NR
Yaa-Arg-Xaa	NR
Ala-Arg	NR
Arg-Arg	NR
Asn-Arg	NR
Asp-Arg	NR
Cys-Arg	NR
Gln-Arg	NR
Glu-Arg	NR
Gly-Arg	NR
His-arg	NR
lle-Arg	NR
Leu-Arg	NR
Lys-Arg	NR
Met-Arg	NR
Phe-Arg	NR
Ser-Arg	NR
Thr-Arg	NR
Trp-Arg	NR
Tyr-Arg	NR
Val-Arg	NR
Ala-Glu-Arg	NR
Arg-Glu-Arg	NR
Asn-Glu-Arg	NR
Asp-Glu-Arg	NR
Cys-Glu-Arg	NR
Gin-Glu-Arg	NR
Giu-Giu-Arg	NR
Gly-Glu-Arg	NR
His-Glu-Arg	NR
lle-Glu-Arg	NR
Leu-Glu-Arg	NR
Lys-Glu-Arg	. NR
Met-Glu-Arg	NR
Phe-Glu-Arg	NR.
Pro-Glu-Arg	NR NR
Ser-Glu-Arg	- NR
Thr-Glu-Arg	NR NR
Trp-Glu-Arg	NR NR
Tyr-Glu-Arg	NR NR
Val-Glu-Arg	NR

	NR
Arg-Ala	NR NR
Arg-Asp	NR NR
Arg-Cys	NR NR
Arg-Gin	NR NR
Arg-Glu	NR NR
Arg-Gly	NR NR
Arg-His	NR NR
Arg-ile	NR NR
Arg-Leu	NR NR
Arg-Lys	NR NR
Arg-Met	NR NR
Arg-Phe	NR NR
Arg-Pro	
Arg-Ser	NR NR
Arg-Thr	NR NR
Arg-Trp	NR
Arg-Tyr	NR
Arg-Val	NR
Arg-Glu-Ala	NR
Arg-Glu-Asn	NR
Arg-Glu-Asp	NR
Arg-Glu-Cys	NR
Arg-Glu-Gln	NR
Arg-Glu-Glu	NR
Arg-Glu-Gly	NR
Arg-Glu-His	NR
Arg-Glu-lle	NR
Arg-Glu-Leu	NR
Arg-Glu-Lys	NR
Arg-Glu-Met	NR
Arg-Glu-Phe	NR
Arg-Glu-Pro	NR
Arg-Glu-Ser	NR
Arg-Glu-Thr	NR
Arg-Glu-Trp	NR
Arg-Glu-Tyr	NR
Arg-Glu-Val	NR
Ala-Arg-Glu	NR
Arg-Arg-Glu	NR
Asn-Arg-Glu	NR
Asp-Arg-Glu	NR
Cys-Arg-Glu	NR
Gin-Arg-Giu	NR
Glu-Arg-Glu	NR
Gly-Arg-Glu	NR
His-Arg-Glu	- NR
Ile-Arg-Glu	NR
Leu-Arg-Glu	NR
Lys-Arg-Glu	NR
Met-Arg-Glu	NR
morring one	

Phe-Arg-Glu	NR NR
Pro-Arg-Glu	NR NR
Ser-Arg-Glu	NR
Thr-Arg-Glu	NR NR
Trp-Arg-Glu	NR
Tyr-Arg-Glu	NR
Val-Arg-Glu	NR NR
Glu-Arg-Ala,	NR
Glu-Arg-Arg	NR
Glu-Arg-Asn	. NR
Glu-Arg-Asp	NR
Glu-Arg-Cys	NR
Glu-Arg-Gin	NR
Glu-Arg-Gly	NR
Glu-Arg-His	. NR
Glu-Arg-ris Glu-Arg-lle	NR
Glu-Arg-lie	NR
Glu-Arg-Les Glu-Arg-Lys	NR
Glu-Arg-Lys Glu-Arg-Met	NR
Glu-Arg-Met	NR
Glu-Arg-Pro	NR
Glu-Arg-Fro	NR
	NR
Glu-Arg-Thr	NR
Glu-Arg-Trp	NR
Glu-Arg-Tyr	NR
Glu-Arg-Val	

Table 20—Additional Exemplary Pharmacologically Active Peptides

Sequence/structure	SEQ ID NO:	Activity
VEPNCDIHVMWEWECFERL	1027	VEGF-antagonist
GERWCFDGPLTWVCGEES	1084	VEGF-antagonist
RGWVEICVADDNGMCVTEAQ	1085	VEGF-antagonist
GWDECDVARMWEWECFAGV	1086	VEGF- antagonist
GERWCFDGPRAWVCGWEI	501	VEGF- antagonist
EELWCFDGPRAWVCGYVK	502	VEGF- antagonist
RGWVEICAADDYGRCLTEAQ	1031	VEGF- antagonist
RGWVEICESDVWGRCL	1087	VEGF- antagonist
RGWVEICESDVWGRCL	1088	VEGF- antagonist
GGNECDIARMWEWECFERL	1089	VEGF- antagonist
RGWVEICAADDYGRCL	1090	VEGF-antagonist
CTTHWGFTLC	1028	MMP inhibitor
CLRSGXGC	1091	MMP inhibitor
CXXHWGFXXC	1092	MMP inhibitor
CXPXC	1093	MMP inhibitor
CRRHWGFEFC	1094	MMP inhibitor
STTHWGFTLS	1095	MMP inhibitor
CSLHWGFWWC	1096	CTLA4-mimetic
GFVCSGIFAVGVGRC	125	CTLA4-mimetic
APGVRLGCAVLGRYC	126	CTLA4-mimetic
LLGRMK	105	Antiviral (HBV)
ICVVQDWGHHRCTAGHMANLTSHASAI	127	C3b antagonist
ICVVQDWGHHRCT	128	C3b antagonist
CVVQDWGHHAC	129	C3b antagonist
STGGFDDVYDWARGVSSALTTTLVATR	185	Vinculin-binding
STGGFDDVYDWARRVSSALTTTLVATR	186	Vinculin-binding
SRGVNFSEWLYDMSAAMKEASNVFPSRRSR	187	Vinculin-binding
SSQNWDMEAGVEDLTAAMLGLLSTIHSSSR	188	Vinculin-binding
SSPSLYTQFLVNYESAATRIQDLLIASRPSR	189	Vinculin-binding
SSTGWVDLLGALQRAADATRTSIPPSLQNSR	190	Vinculin-binding
DVYTKKELIECARRVSEK	191	Vinculin-binding
EKGSYYPGSGIAQFHIDYNNVS	192	C4BP-binding
SGIAQFHIDYNNVSSAEGWHVN	193	C4BP-binding
LVTVEKGSYYPGSGIAQFHIDYNNVSSAEGWHVN	194	C4BP-binding
SGIAQFHIDYNNVS	195	C4BP-binding
LLGRMK	279	anti-HBV
ALLGRMKG	280	anti-HBV
LDPAFR	281	anti-HBV
CXXRGDC	322	Inhibition of platelet
		aggregation
RPLPPLP	323	Src antagonist
PPVPPR	324	Src antagonist
XFXDXWXXLXX	325	Anti-cancer
	1	(particularly for

		sarcomas)
KACRRLFGPVDSEQLSRDCD	326	p16-mimetic
RERWNFDFVTETPLEGDFAW	327	p16-mimetic
KRRQTSMTDFYHSKRRLIFS	328	p16-mimetic
TSMTDFYHSKRRLIFSKRKP	329	p16-mimetic
RRLIF	330	p16-mimetic
KRRQTSATDFYHSKRRLIFSRQIKIWFQNRRMKWKK	331	p16-mimetic
KRRLIFSKRQIKIWFQNRRMKWKK	332	p16-mimetic
Asn Gin Gly Arg His Phe Cys Gly Gly Ala Leu lie His Ala	498	CAP37 mimetic/LPS
Arg Phe Val Met Thr Ala Ala Ser Cys Phe Gin		binding
Arg His Phe Cys Gly Gly Ala Leu lle His Ala Arg Phe Val	499	CAP37 mimetic/LPS
Met Thr Ala Ala Ser Cvs		binding
Gly Thr Arg Cys Gln Val Ala Gly Trp Gly Ser Gln Arg Ser	500	CAP37 mimetic/LPS
Gly Gly Arg Leu Ser Arg Phe Pro Arg Phe Val Asn Val		binding
WHWRHRIPLQLAAGR	1097	carbohydrate (GD1
		alpha) mimetic
	1098	β2GPI Ab binding
LKTPRV	1099	β2GPI Ab binding
NTLKTPRV		
NTLKTPRVGGC	1100	β2GPI Ab binding
KDKATF	1101	β2GPI Ab binding
KDKATFGCHD	1102	β2GPI Ab binding
KDKATFGCHDGC	1103	β2GPI Ab binding
TLRVYK	1104	β2GPI Ab binding
ATLRVYKGG	1105	β2GPI Ab binding
CATLRVYKGG	1106	β2GPI Ab binding
INLKALAALAKKIL	1107	Membrane-
		transporting
GWT	NR	Membrane-
·	1	transporting
GWTLNSAGYLLG	1108	Membrane-
	1100	transporting
GWTLNSAGYLLGKINLKALAALAKKIL	1109	Membrane-
	 _	transporting

The present invention is also particularly useful with peptides having activity in treatment of:

 cancer, wherein the peptide is a VEGF-mimetic or a VEGF receptor antagonist, a HER2 agonist or antagonist, a CD20 antagonist and the like;

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- asthma, wherein the protein of interest is a CKR3 antagonist, an IL-5 receptor antagonist, and the like;
- thrombosis, wherein the protein of interest is a GPIIb antagonist, a GPIIIa antagonist, and the like;

 autoimmune diseases and other conditions involving immune modulation, wherein the protein of interest is an IL-2 receptor antagonist, a CD40 agonist or antagonist, a CD40L agonist or antagonist, a thymopoietin mimetic and the like.

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<u>Vehicles</u>. This invention requires the presence of at least one vehicle (F¹, F²) attached to a peptide through the N-terminus, C-terminus or a sidechain of one of the amino acid residues. Multiple vehicles may also be used; e.g., Fc's at each terminus or an Fc at a terminus and a PEG group at the other terminus or a sidechain.

An Fc domain is the preferred vehicle. The Fc domain may be fused to the N or C termini of the peptides or at both the N and C termini. For the TPO-mimetic peptides, molecules having the Fc domain fused to the N terminus of the peptide portion of the molecule are more bioactive than other such fusions, so fusion to the N terminus is preferred.

As noted above, Fc variants are suitable vehicles within the scope of this invention. A native Fc may be extensively modified to form an Fc variant in accordance with this invention, provided binding to the salvage receptor is maintained; see, for example WO 97/34631 and WO 96/32478. In such Fc variants, one may remove one or more sites of a native Fc that provide structural features or functional activity not required by the fusion molecules of this invention. One may remove these sites by, for example, substituting or deleting residues, inserting residues into the site, or truncating portions containing the site. The inserted or substituted residues may also be altered amino acids, such as peptidomimetics or D-amino acids. Fc variants may be desirable for a number of reasons, several of which are described below. Exemplary Fc variants include molecules and sequences in which:

1. Sites involved in disulfide bond formation are removed. Such removal may avoid reaction with other cysteine-containing proteins present in

the host cell used to produce the molecules of the invention. For this purpose, the cysteine-containing segment at the N-terminus may be truncated or cysteine residues may be deleted or substituted with other amino acids (e.g., alanyl, seryl). In particular, one may truncate the N-terminal 20-amino acid segment of SEQ ID NO: 2 or delete or substitute the cysteine residues at positions 7 and 10 of SEQ ID NO: 2. Even when cysteine residues are removed, the single chain Fc domains can still form a dimeric Fc domain that is held together non-covalently.

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- 2. A native Fc is modified to make it more compatible with a selected host cell. For example, one may remove the PA sequence near the N-terminus of a typical native Fc, which may be recognized by a digestive enzyme in <u>E. coli</u> such as proline iminopeptidase. One may also add an N-terminal methionine residue, especially when the molecule is expressed recombinantly in a bacterial cell such as <u>E. coli</u>. The Fc domain of SEQ ID NO: 2 (Figure 4) is one such Fc variant.
 - 3. A portion of the N-terminus of a native Fc is removed to prevent N-terminal heterogeneity when expressed in a selected host cell. For this purpose, one may delete any of the first 20 amino acid residues at the N-terminus, particularly those at positions 1, 2, 3, 4 and 5.
- 4. One or more glycosylation sites are removed. Residues that are typically glycosylated (e.g., asparagine) may confer cytolytic response. Such residues may be deleted or substituted with unglycosylated residues (e.g., alanine).
- 5. Sites involved in interaction with complement, such as the C1q binding site, are removed. For example, one may delete or substitute the EKK sequence of human IgG1. Complement recruitment may not be advantageous for the molecules of this invention and so may be avoided with such an Fc variant.

6. Sites are removed that affect binding to Fc receptors other than a salvage receptor. A native Fc may have sites for interaction with certain white blood cells that are not required for the fusion molecules of the present invention and so may be removed.

- 7. The ADCC site is removed. ADCC sites are known in the art; see, for example, <u>Molec. Immunol</u>. 29 (5): 633-9 (1992) with regard to ADCC sites in IgG1. These sites, as well, are not required for the fusion molecules of the present invention and so may be removed.
- 8. When the native Fc is derived from a non-human antibody, the native Fc may be humanized. Typically, to humanize a native Fc, one will substitute selected residues in the non-human native Fc with residues that are normally found in human native Fc. Techniques for antibody humanization are well known in the art.

Preferred Fc variants include the following. In SEQ ID NO: 2

(Figure 4) the leucine at position 15 may be substituted with glutamate; the glutamate at position 99, with alanine; and the lysines at positions 101 and 103, with alanines. In addition, one or more tyrosine residues can be replaced by phenyalanine residues.

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An alternative vehicle would be a protein, polypeptide, peptide, antibody, antibody fragment, , or small molecule (e.g., a peptidomimetic compound) capable of binding to a salvage receptor. For example, one could use as a vehicle a polypeptide as described in U.S. Pat. No. 5,739,277, issued April 14, 1998 to Presta et al. Peptides could also be selected by phage display for binding to the FcRn salvage receptor. Such salvage receptor-binding compounds are also included within the meaning of "vehicle" and are within the scope of this invention. Such vehicles should be selected for increased half-life (e.g., by avoiding sequences recognized by proteases) and decreased immunogenicity (e.g., by favoring non-immunogenic sequences, as discovered in antibody humanization).

As noted above, polymer vehicles may also be used for F¹ and F². Various means for attaching chemical moieties useful as vehicles are currently available, see, e.g., Patent Cooperation Treaty ("PCT") International Publication No. WO 96/11953, entitled "N-Terminally Chemically Modified Protein Compositions and Methods," herein incorporated by reference in its entirety. This PCT publication discloses, among other things, the selective attachment of water soluble polymers to the N-terminus of proteins.

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A preferred polymer vehicle is polyethylene glycol (PEG). The PEG group may be of any convenient molecular weight and may be linear or branched. The average molecular weight of the PEG will preferably range from about 2 kiloDalton ("kD") to about 100 kDa, more preferably from about 5 kDa to about 50 kDa, most preferably from about 5 kDa to about 10 kDa. The PEG groups will generally be attached to the compounds of the invention via acylation or reductive alkylation through a reactive group on the PEG moiety (e.g., an aldehyde, amino, thiol, or ester group) to a reactive group on the inventive compound (e.g., an aldehyde, amino, or ester group).

A useful strategy for the PEGylation of synthetic peptides consists of combining, through forming a conjugate linkage in solution, a peptide and a PEG moiety, each bearing a special functionality that is mutually reactive toward the other. The peptides can be easily prepared with conventional solid phase synthesis (see, for example, Figures 5 and 6 and the accompanying text herein). The peptides are "preactivated" with an appropriate functional group at a specific site. The precursors are purified and fully characterized prior to reacting with the PEG moiety. Ligation of the peptide with PEG usually takes place in aqueous phase and can be easily monitored by reverse phase analytical HPLC. The PEGylated peptides can be easily purified by preparative HPLC and characterized by

analytical HPLC, amino acid analysis and laser desorption mass spectrometry.

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Polysaccharide polymers are another type of water soluble polymer which may be used for protein modification. Dextrans are polysaccharide polymers comprised of individual subunits of glucose predominantly linked by $\alpha 1$ -6 linkages. The dextran itself is available in many molecular weight ranges, and is readily available in molecular weights from about 1 kD to about 70 kD. Dextran is a suitable water soluble polymer for use in the present invention as a vehicle by itself or in combination with another vehicle (e.g., Fc). See, for example, WO 96/11953 and WO 96/05309. The use of dextran conjugated to therapeutic or diagnostic immunoglobulins has been reported; see, for example, European Patent Publication No. 0 315 456, which is hereby incorporated by reference. Dextran of about 1 kD to about 20 kD is preferred when dextran is used as a vehicle in accordance with the present invention.

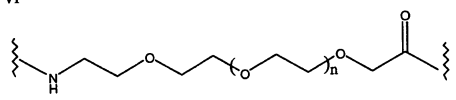
Linkers. Any "linker" group is optional. When present, its chemical structure is not critical, since it serves primarily as a spacer. The linker is preferably made up of amino acids linked together by peptide bonds. Thus, in preferred embodiments, the linker is made up of from 1 to 20 amino acids linked by peptide bonds, wherein the amino acids are selected from the 20 naturally occurring amino acids. Some of these amino acids may be glycosylated, as is well understood by those in the art. In a more preferred embodiment, the 1 to 20 amino acids are selected from glycine, alanine, proline, asparagine, glutamine, and lysine. Even more preferably, a linker is made up of a majority of amino acids that are sterically unhindered, such as glycine and alanine. Thus, preferred linkers are polyglycines (particularly (Gly), (Gly), poly(Gly-Ala), and polyalanines. Other specific examples of linkers are:

(Gly)₃Lys(Gly)₄ (SEQ ID NO: 333);

(Gly)₃AsnGlySer(Gly)₂ (SEQ ID NO: 334); (Gly)₃Cys(Gly)₄ (SEQ ID NO: 335); and GlyProAsnGlyGly (SEQ ID NO: 336).

To explain the above nomenclature, for example, (Gly)₃Lys(Gly)₄ means Gly-Gly-Gly-Gly-Gly-Gly-Gly. Combinations of Gly and Ala are also preferred. The linkers shown here are exemplary; linkers within the scope of this invention may be much longer and may include other residues.

Non-peptide linkers are also possible. For example, alkyl linkers such as -NH-(CH_2), -C(O)-, wherein s = 2-20 could be used. These alkyl linkers may further be substituted by any non-sterically hindering group such as lower alkyl (e.g., C_1 - C_6) lower acyl, halogen (e.g., Cl, Br), CN, NH₂, phenyl, etc. An exemplary non-peptide linker is a PEG linker, VI



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wherein n is such that the linker has a molecular weight of 100 to 5000 kD, preferably 100 to 500 kD. The peptide linkers may be altered to form derivatives in the same manner as described above.

Derivatives. The inventors also contemplate derivatizing the
peptide and/or vehicle portion of the compounds. Such derivatives may
improve the solubility, absorption, biological half life, and the like of the
compounds. The moieties may alternatively eliminate or attenuate any
undesirable side-effect of the compounds and the like. Exemplary
derivatives include compounds in which:

The compound or some portion thereof is cyclic. For example, the
peptide portion may be modified to contain two or more Cys residues
(e.g., in the linker), which could cyclize by disulfide bond formation.

For citations to references on preparation of cyclized derivatives, see Table 2.

2. The compound is cross-linked or is rendered capable of cross-linking between molecules. For example, the peptide portion may be modified to contain one Cys residue and thereby be able to form an intermolecular disulfide bond with a like molecule. The compound may also be cross-linked through its C-terminus, as in the molecule shown below.

VII

$$F^{1}-(X^{1})_{b}-CO-N$$
 NH_{2}
 NH_{2}

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- 4. One or more peptidyl [-C(O)NR-] linkages (bonds) is replaced by a non-peptidyl linkage. Exemplary non-peptidyl linkages are -CH₂-carbamate [-CH₂-OC(O)NR-], phosphonate , -CH₂-sulfonamide [-CH₂-S(O)₂NR-], urea [-NHC(O)NH-], -CH₂-secondary amine, and alkylated peptide [-C(O)NR⁶- wherein R⁶ is lower alkyl].
- 5. The N-terminus is derivatized. Typically, the N-terminus may be acylated or modified to a substituted amine. Exemplary N-terminal derivative groups include -NRR¹ (other than -NH₂), -NRC(O)R¹, -NRC(O)OR¹, -NRS(O)₂R¹, -NHC(O)NHR¹, succinimide, or benzyloxycarbonyl-NH- (CBZ-NH-), wherein R and R¹ are each independently hydrogen or lower alkyl and wherein the phenyl ring may be substituted with 1 to 3 substituents selected from the group consisting of C₁-C₄ alkyl, C₁-C₄ alkoxy, chloro, and bromo.
- 6. The free C-terminus is derivatized. Typically, the C-terminus is esterified or amidated. For example, one may use methods described in the art to add (NH-CH₂-CH₂-NH₂)₂ to compounds of this invention

having any of SEQ ID NOS: 504 to 508 at the C-terminus. Likewise, one may use methods described in the art to add -NH₂ to compounds of this invention having any of SEQ ID NOS: 924 to 955, 963 to 972, 1005 to 1013, or 1018 to 1023 at the C-terminus. Exemplary C-terminal derivative groups include, for example, -C(O)R² wherein R² is lower alkoxy or -NR³R⁴ wherein R³ and R⁴ are independently hydrogen or C₁-C₈ alkyl (preferably C₁-C₄ alkyl).

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- A disulfide bond is replaced with another, preferably more stable, cross-linking moiety (e.g., an alkylene). See, e.g., Bhatnagar et al. (1996), J. Med. Chem. 39: 3814-9; Alberts et al. (1993) Thirteenth Am. Pep. Symp., 357-9.
- 8. One or more individual amino acid residues is modified. Various derivatizing agents are known to react specifically with selected sidechains or terminal residues, as described in detail below.

Lysinyl residues and amino terminal residues may be reacted with succinic or other carboxylic acid anhydrides, which reverse the charge of the lysinyl residues. Other suitable reagents for derivatizing alpha-amino-containing residues include imidoesters such as methyl picolinimidate; pyridoxal phosphate; pyridoxal; chloroborohydride; trinitrobenzenesulfonic acid; O-methylisourea; 2,4 pentanedione; and transaminase-catalyzed reaction with glyoxylate.

Arginyl residues may be modified by reaction with any one or combination of several conventional reagents, including phenylglyoxal, 2,3-butanedione, 1,2-cyclohexanedione, and ninhydrin. Derivatization of arginyl residues requires that the reaction be performed in alkaline conditions because of the high pKa of the guanidine functional group. Furthermore, these reagents may react with the groups of lysine as well as the arginine epsilon-amino group.

Specific modification of tyrosyl residues has been studied extensively, with particular interest in introducing spectral labels into tyrosyl residues by reaction with aromatic diazonium compounds or tetranitromethane. Most commonly, N-acetylimidizole and tetranitromethane are used to form O-acetyl tyrosyl species and 3-nitro derivatives, respectively.

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Carboxyl sidechain groups (aspartyl or glutamyl) may be selectively modified by reaction with carbodiimides (R'-N=C=N-R') such as 1-cyclohexyl-3-(2-morpholinyl-(4-ethyl) carbodiimide or 1-ethyl-3-(4-azonia-4,4-dimethylpentyl) carbodiimide. Furthermore, aspartyl and glutamyl residues may be converted to asparaginyl and glutaminyl residues by reaction with ammonium ions.

Glutaminyl and asparaginyl residues may be deamidated to the corresponding glutamyl and aspartyl residues. Alternatively, these residues are deamidated under mildly acidic conditions. Either form of these residues falls within the scope of this invention.

Cysteinyl residues can be replaced by amino acid residues or other moieties either to eliminate disulfide bonding or, conversely, to stabilize cross-linking. See, e.g., Bhatnagar <u>et al.</u> (1996), <u>J. Med. Chem.</u> 39: 3814-9.

Derivatization with bifunctional agents is useful for cross-linking the peptides or their functional derivatives to a water-insoluble support matrix or to other macromolecular vehicles. Commonly used cross-linking agents include, e.g., 1,1-bis(diazoacetyl)-2-phenylethane, glutaraldehyde, N-hydroxysuccinimide esters, for example, esters with 4-azidosalicylic acid, homobifunctional imidoesters, including disuccinimidyl esters such as 3,3'-dithiobis(succinimidylpropionate), and bifunctional maleimides such as bis-N-maleimido-1,8-octane. Derivatizing agents such as methyl-3-[(p-azidophenyl)dithio]propioimidate yield photoactivatable intermediates that are capable of forming crosslinks in the presence of light. Alternatively, reactive water-insoluble matrices such as cyanogen bromide-activated carbohydrates

and the reactive substrates described in U.S. Pat. Nos. 3,969,287; 3,691,016; 4,195,128; 4,247,642; 4,229,537; and 4,330,440 are employed for protein immobilization.

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Carbohydrate (oligosaccharide) groups may conveniently be attached to sites that are known to be glycosylation sites in proteins. Generally, O-linked oligosaccharides are attached to serine (Ser) or threonine (Thr) residues while N-linked oligosaccharides are attached to asparagine (Asn) residues when they are part of the sequence Asn-X-Ser/Thr, where X can be any amino acid except proline. X is preferably one of the 19 naturally occurring amino acids other than proline. The structures of N-linked and O-linked oligosaccharides and the sugar residues found in each type are different. One type of sugar that is commonly found on both is N-acetylneuraminic acid (referred to as sialic acid). Sialic acid is usually the terminal residue of both N-linked and Olinked oligosaccharides and, by virtue of its negative charge, may confer acidic properties to the glycosylated compound. Such site(s) may be incorporated in the linker of the compounds of this invention and are preferably glycosylated by a cell during recombinant production of the polypeptide compounds (e.g., in mammalian cells such as CHO, BHK, COS). However, such sites may further be glycosylated by synthetic or semi-synthetic procedures known in the art.

Other possible modifications include hydroxylation of proline and lysine, phosphorylation of hydroxyl groups of seryl or threonyl residues, oxidation of the sulfur atom in Cys, methylation of the alpha-amino groups of lysine, arginine, and histidine side chains. Creighton, <u>Proteins: Structure and Molecule Properties</u> (W. H. Freeman & Co., San Francisco), pp. 79-86 (1983).

Compounds of the present invention may be changed at the DNA level, as well. The DNA sequence of any portion of the compound may be

changed to codons more compatible with the chosen host cell. For <u>E. coli</u>, which is the preferred host cell, optimized codons are known in the art. Codons may be substituted to eliminate restriction sites or to include silent restriction sites, which may aid in processing of the DNA in the selected host cell. The vehicle, linker and peptide DNA sequences may be modified to include any of the foregoing sequence changes.

Methods of Making

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The compounds of this invention largely may be made in transformed host cells using recombinant DNA techniques. To do so, a recombinant DNA molecule coding for the peptide is prepared. Methods of preparing such DNA molecules are well known in the art. For instance, sequences coding for the peptides could be excised from DNA using suitable restriction enzymes. Alternatively, the DNA molecule could be synthesized using chemical synthesis techniques, such as the phosphoramidate method. Also, a combination of these techniques could be used.

The invention also includes a vector capable of expressing the peptides in an appropriate host. The vector comprises the DNA molecule that codes for the peptides operatively linked to appropriate expression control sequences. Methods of effecting this operative linking, either before or after the DNA molecule is inserted into the vector, are well known. Expression control sequences include promoters, activators, enhancers, operators, ribosomal binding sites, start signals, stop signals, cap signals, polyadenylation signals, and other signals involved with the control of transcription or translation.

The resulting vector having the DNA molecule thereon is used to transform an appropriate host. This transformation may be performed using methods well known in the art.

Any of a large number of available and well-known host cells may be used in the practice of this invention. The selection of a particular host is dependent upon a number of factors recognized by the art. These include, for example, compatibility with the chosen expression vector, toxicity of the peptides encoded by the DNA molecule, rate of transformation, ease of recovery of the peptides, expression characteristics, bio-safety and costs. A balance of these factors must be struck with the understanding that not all hosts may be equally effective for the expression of a particular DNA sequence. Within these general guidelines, useful microbial hosts include bacteria (such as <u>E. coli</u> sp.), yeast (such as <u>Saccharomyces</u> sp.) and other fungi, insects, plants, mammalian (including human) cells in culture, or other hosts known in the art.

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Next, the transformed host is cultured and purified. Host cells may be cultured under conventional fermentation conditions so that the desired compounds are expressed. Such fermentation conditions are well known in the art. Finally, the peptides are purified from culture by methods well known in the art.

The compounds may also be made by synthetic methods. For example, solid phase synthesis techniques may be used. Suitable techniques are well known in the art, and include those described in Merrifield (1973), Chem. Polypeptides, pp. 335-61 (Katsoyannis and Panayotis eds.); Merrifield (1963), J. Am. Chem. Soc. 85: 2149; Davis et al. (1985), Biochem. Intl. 10: 394-414; Stewart and Young (1969), Solid Phase Peptide Synthesis; U.S. Pat. No. 3,941,763; Finn et al. (1976), The Proteins (3rd ed.) 2: 105-253; and Erickson et al. (1976), The Proteins (3rd ed.) 2: 257-527. Solid phase synthesis is the preferred technique of making individual peptides since it is the most cost-effective method of making small peptides.

Compounds that contain derivatized peptides or which contain non-peptide groups may be synthesized by well-known organic chemistry techniques.

Uses of the Compounds

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<u>In general</u>. The compounds of this invention have pharmacologic activity resulting from their ability to bind to proteins of interest as agonists, mimetics or antagonists of the native ligands of such proteins of interest. The utility of specific compounds is shown in Table 2. The activity of these compounds can be measured by assays known in the art. For the TPO-mimetic and EPO-mimetic compounds, <u>in vivo</u> assays are further described in the Examples section herein.

In addition to therapeutic uses, the compounds of the present invention are useful in diagnosing diseases characterized by dysfunction of their associated protein of interest. In one embodiment, a method of detecting in a biological sample a protein of interest (e.g., a receptor) that is capable of being activated comprising the steps of: (a) contacting the sample with a compound of this invention; and (b) detecting activation of the protein of interest by the compound. The biological samples include tissue specimens, intact cells, or extracts thereof. The compounds of this invention may be used as part of a diagnostic kit to detect the presence of their associated proteins of interest in a biological sample. Such kits employ the compounds of the invention having an attached label to allow for detection. The compounds are useful for identifying normal or abnormal proteins of interest. For the EPO-mimetic compounds, for example, presence of abnormal protein of interest in a biological sample may be indicative of such disorders as Diamond Blackfan anemia, where it is believed that the EPO receptor is dysfunctional.

Therapeutic uses of EPO-mimetic compounds. The EPO-mimetic compounds of the invention are useful for treating disorders characterized by low red blood cell levels. Included in the invention are methods of modulating the endogenous activity of an EPO receptor in a mammal, preferably methods of increasing the activity of an EPO receptor. In

general, any condition treatable by erythropoietin, such as anemia, may also be treated by the EPO-mimetic compounds of the invention. These compounds are administered by an amount and route of delivery that is appropriate for the nature and severity of the condition being treated and may be ascertained by one skilled in the art. Preferably, administration is by injection, either subcutaneous, intramuscular, or intravenous.

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Therapeutic uses of TPO-mimetic compounds. For the TPO-mimetic compounds, one can utilize such standard assays as those described in WO95/26746 entitled "Compositions and Methods for Stimulating Megakaryocyte Growth and Differentiation". In vivo assays also appear in the Examples hereinafter.

The conditions to be treated are generally those that involve an existing megakaryocyte/platelet deficiency or an expected megakaryocyte/platelet deficiency (e.g., because of planned surgery or platelet donation). Such conditions will usually be the result of a deficiency (temporary or permanent) of active Mpl ligand in vivo. The generic term for platelet deficiency is thrombocytopenia, and hence the methods and compositions of the present invention are generally available for treating thrombocytopenia in patients in need thereof.

Thrombocytopenia (platelet deficiencies) may be present for various reasons, including chemotherapy and other therapy with a variety of drugs, radiation therapy, surgery, accidental blood loss, and other specific disease conditions. Exemplary specific disease conditions that involve thrombocytopenia and may be treated in accordance with this invention are: aplastic anemia, idiopathic thrombocytopenia, metastatic tumors which result in thrombocytopenia, systemic lupus erythematosus, splenomegaly, Fanconi's syndrome, vitamin B12 deficiency, folic acid deficiency, May-Hegglin anomaly, Wiskott-Aldrich syndrome, and paroxysmal nocturnal hemoglobinuria. Also, certain treatments for AIDS

result in thrombocytopenia (e.g., AZT). Certain wound healing disorders might also benefit from an increase in platelet numbers.

With regard to anticipated platelet deficiencies, e.g., due to future surgery, a compound of the present invention could be administered several days to several hours prior to the need for platelets. With regard to acute situations, e.g., accidental and massive blood loss, a compound of this invention could be administered along with blood or purified platelets.

The TPO-mimetic compounds of this invention may also be useful in stimulating certain cell types other than megakaryocytes if such cells are found to express Mpl receptor. Conditions associated with such cells that express the Mpl receptor, which are responsive to stimulation by the Mpl ligand, are also within the scope of this invention.

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The TPO-mimetic compounds of this invention may be used in any situation in which production of platelets or platelet precursor cells is desired, or in which stimulation of the c-Mpl receptor is desired. Thus, for example, the compounds of this invention may be used to treat any condition in a mammal wherein there is a need of platelets, megakaryocytes, and the like. Such conditions are described in detail in the following exemplary sources: WO95/26746; WO95/21919; WO95/18858; WO95/21920 and are incorporated herein.

The TPO-mimetic compounds of this invention may also be useful in maintaining the viability or storage life of platelets and/or megakaryocytes and related cells. Accordingly, it could be useful to include an effective amount of one or more such compounds in a composition containing such cells.

The therapeutic methods, compositions and compounds of the present invention may also be employed, alone or in combination with other cytokines, soluble Mpl receptor, hematopoietic factors, interleukins, growth factors or antibodies in the treatment of disease states

characterized by other symptoms as well as platelet deficiencies. It is anticipated that the inventive compound will prove useful in treating some forms of thrombocytopenia in combination with general stimulators of hematopoiesis, such as IL-3 or GM-CSF. Other megakaryocytic stimulatory factors, i.e., meg-CSF, stem cell factor (SCF), leukemia 5 inhibitory factor (LIF), oncostatin M (OSM), or other molecules with megakaryocyte stimulating activity may also be employed with Mpl ligand. Additional exemplary cytokines or hematopoietic factors for such co-administration include IL-1 alpha, IL-1 beta, IL-2, IL-3, IL-4, IL-5, IL-6, IL-11, colony stimulating factor-1 (CSF-1), SCF, GM-CSF, granulocyte colony stimulating factor (G-CSF), EPO, interferon-alpha (IFN-alpha), consensus interferon, IFN-beta, or IFN-gamma. It may further be useful to administer, either simultaneously or sequentially, an effective amount of a soluble mammalian Mpl receptor, which appears to have an effect of causing megakaryocytes to fragment into platelets once the megakaryocytes have reached mature form. Thus, administration of an inventive compound (to enhance the number of mature megakaryocytes) followed by administration of the soluble Mpl receptor (to inactivate the ligand and allow the mature megakaryocytes to produce platelets) is expected to be a particularly effective means of stimulating platelet production. The dosage recited above would be adjusted to compensate for such additional components in the therapeutic composition. Progress of the treated patient can be monitored by conventional methods.

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In cases where the inventive compounds are added to compositions of platelets and/or megakaryocytes and related cells, the amount to be included will generally be ascertained experimentally by techniques and assays known in the art. An exemplary range of amounts is 0.1 µg—1 mg inventive compound per 10° cells.

Pharmaceutical Compositions

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In General. The present invention also provides methods of using pharmaceutical compositions of the inventive compounds. Such pharmaceutical compositions may be for administration for injection, or for oral, pulmonary, nasal, transdermal or other forms of administration. In general, the invention encompasses pharmaceutical compositions comprising effective amounts of a compound of the invention together with pharmaceutically acceptable diluents, preservatives, solubilizers, emulsifiers, adjuvants and/or carriers. Such compositions include diluents of various buffer content (e.g., Tris-HCl, acetate, phosphate), pH and ionic strength; additives such as detergents and solubilizing agents (e.g., Tween 80, Polysorbate 80), anti-oxidants (e.g., ascorbic acid, sodium metabisulfite), preservatives (e.g., Thimersol, benzyl alcohol) and bulking substances (e.g., lactose, mannitol); incorporation of the material into particulate preparations of polymeric compounds such as polylactic acid, polyglycolic acid, etc. or into liposomes. Hyaluronic acid may also be used, and this may have the effect of promoting sustained duration in the circulation. Such compositions may influence the physical state, stability, rate of in vivo release, and rate of in vivo clearance of the present proteins and derivatives. See, e.g., Remington's Pharmaceutical Sciences, 18th Ed. (1990, Mack Publishing Co., Easton, PA 18042) pages 1435-1712 which are herein incorporated by reference. The compositions may be prepared in liquid form, or may be in dried powder, such as lyophilized form. Implantable sustained release formulations are also contemplated, as are transdermal formulations.

Oral dosage forms. Contemplated for use herein are oral solid dosage forms, which are described generally in Chapter 89 of Remington's Pharmaceutical Sciences (1990), 18th Ed., Mack Publishing Co. Easton PA 18042, which is herein incorporated by reference. Solid dosage forms include tablets, capsules, pills, troches or lozenges, cachets or pellets. Also,

liposomal or proteinoid encapsulation may be used to formulate the present compositions (as, for example, proteinoid microspheres reported in U.S. Patent No. 4,925,673). Liposomal encapsulation may be used and the liposomes may be derivatized with various polymers (e.g., U.S. Patent No. 5,013,556). A description of possible solid dosage forms for the therapeutic is given in Chapter 10 of Marshall, K., Modern Pharmaceutics (1979), edited by G. S. Banker and C. T. Rhodes, herein incorporated by reference. In general, the formulation will include the inventive compound, and inert ingredients which allow for protection against the stomach environment, and release of the biologically active material in the intestine.

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Also specifically contemplated are oral dosage forms of the above inventive compounds. If necessary, the compounds may be chemically modified so that oral delivery is efficacious. Generally, the chemical modification contemplated is the attachment of at least one moiety to the compound molecule itself, where said moiety permits (a) inhibition of proteolysis; and (b) uptake into the blood stream from the stomach or intestine. Also desired is the increase in overall stability of the compound and increase in circulation time in the body. Moieties useful as covalently attached vehicles in this invention may also be used for this purpose. Examples of such moieties include: PEG, copolymers of ethylene glycol and propylene glycol, carboxymethyl cellulose, dextran, polyvinyl alcohol, polyvinyl pyrrolidone and polyproline. See, for example, Abuchowski and Davis, Soluble Polymer-Enzyme Adducts, Enzymes as Drugs (1981), Hocenberg and Roberts, eds., Wiley-Interscience, New York, NY,, pp 367-83; Newmark, et al. (1982), J. Appl. Biochem. 4:185-9. Other polymers that could be used are poly-1,3-dioxolane and poly-1,3,6-tioxocane. Preferred for pharmaceutical usage, as indicated above, are PEG moieties.

For oral delivery dosage forms, it is also possible to use a salt of a modified aliphatic amino acid, such as sodium N-(8-[2-hydroxybenzoyl] amino) caprylate (SNAC), as a carrier to enhance absorption of the therapeutic compounds of this invention. The clinical efficacy of a heparin formulation using SNAC has been demonstrated in a Phase II trial conducted by Emisphere Technologies. See US Patent No. 5,792,451, "Oral drug delivery composition and methods".

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The compounds of this invention can be included in the formulation as fine multiparticulates in the form of granules or pellets of particle size about 1 mm. The formulation of the material for capsule administration could also be as a powder, lightly compressed plugs or even as tablets. The therapeutic could be prepared by compression.

Colorants and flavoring agents may all be included. For example, the protein (or derivative) may be formulated (such as by liposome or microsphere encapsulation) and then further contained within an edible product, such as a refrigerated beverage containing colorants and flavoring agents.

One may dilute or increase the volume of the compound of the invention with an inert material. These diluents could include carbohydrates, especially mannitol, α -lactose, anhydrous lactose, cellulose, sucrose, modified dextrans and starch. Certain inorganic salts may also be used as fillers including calcium triphosphate, magnesium carbonate and sodium chloride. Some commercially available diluents are Fast-Flo, Emdex, STA-Rx 1500, Emcompress and Avicell.

Disintegrants may be included in the formulation of the therapeutic into a solid dosage form. Materials used as disintegrants include but are not limited to starch including the commercial disintegrant based on starch, Explotab. Sodium starch glycolate, Amberlite, sodium carboxymethylcellulose, ultramylopectin, sodium alginate, gelatin, orange

peel, acid carboxymethyl cellulose, natural sponge and bentonite may all be used. Another form of the disintegrants are the insoluble cationic exchange resins. Powdered gums may be used as disintegrants and as binders and these can include powdered gums such as agar, Karaya or tragacanth. Alginic acid and its sodium salt are also useful as disintegrants.

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Binders may be used to hold the therapeutic agent together to form a hard tablet and include materials from natural products such as acacia, tragacanth, starch and gelatin. Others include methyl cellulose (MC), ethyl cellulose (EC) and carboxymethyl cellulose (CMC). Polyvinyl pyrrolidone (PVP) and hydroxypropylmethyl cellulose (HPMC) could both be used in alcoholic solutions to granulate the therapeutic.

An antifrictional agent may be included in the formulation of the therapeutic to prevent sticking during the formulation process. Lubricants may be used as a layer between the therapeutic and the die wall, and these can include but are not limited to; stearic acid including its magnesium and calcium salts, polytetrafluoroethylene (PTFE), liquid paraffin, vegetable oils and waxes. Soluble lubricants may also be used such as sodium lauryl sulfate, magnesium lauryl sulfate, polyethylene glycol of various molecular weights, Carbowax 4000 and 6000.

Glidants that might improve the flow properties of the drug during formulation and to aid rearrangement during compression might be added. The glidants may include starch, talc, pyrogenic silica and hydrated silicoaluminate.

To aid dissolution of the compound of this invention into the aqueous environment a surfactant might be added as a wetting agent. Surfactants may include anionic detergents such as sodium lauryl sulfate, dioctyl sodium sulfosuccinate and dioctyl sodium sulfonate. Cationic detergents might be used and could include benzalkonium chloride or

benzethonium chloride. The list of potential nonionic detergents that could be included in the formulation as surfactants are lauromacrogol 400, polyoxyl 40 stearate, polyoxyethylene hydrogenated castor oil 10, 50 and 60, glycerol monostearate, polysorbate 40, 60, 65 and 80, sucrose fatty acid ester, methyl cellulose and carboxymethyl cellulose. These surfactants could be present in the formulation of the protein or derivative either alone or as a mixture in different ratios.

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Additives may also be included in the formulation to enhance uptake of the compound. Additives potentially having this property are for instance the fatty acids oleic acid, linoleic acid and linolenic acid.

Controlled release formulation may be desirable. The compound of this invention could be incorporated into an inert matrix which permits release by either diffusion or leaching mechanisms e.g., gums. Slowly degenerating matrices may also be incorporated into the formulation, e.g., alginates, polysaccharides. Another form of a controlled release of the compounds of this invention is by a method based on the Oros therapeutic system (Alza Corp.), i.e., the drug is enclosed in a semipermeable membrane which allows water to enter and push drug out through a single small opening due to osmotic effects. Some enteric coatings also have a delayed release effect.

Other coatings may be used for the formulation. These include a variety of sugars which could be applied in a coating pan. The therapeutic agent could also be given in a film coated tablet and the materials used in this instance are divided into 2 groups. The first are the nonenteric materials and include methyl cellulose, ethyl cellulose, hydroxyethyl cellulose, methylhydroxy-ethyl cellulose, hydroxypropyl cellulose, hydroxypropyl-methyl cellulose, sodium carboxy-methyl cellulose, providone and the polyethylene glycols. The second group consists of the enteric materials that are commonly esters of phthalic acid.

A mix of materials might be used to provide the optimum film coating. Film coating may be carried out in a pan coater or in a fluidized bed or by compression coating.

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Pulmonary delivery forms. Also contemplated herein is pulmonary delivery of the present protein (or derivatives thereof). The protein (or derivative) is delivered to the lungs of a mammal while inhaling and traverses across the lung epithelial lining to the blood stream. (Other reports of this include Adjei et al., Pharma. Res. (1990) 7: 565-9; Adjei et al. (1990), Internatl. J. Pharmaceutics 63: 135-44 (leuprolide acetate); Braquet et al. (1989), J. Cardiovasc. Pharmacol. 13 (suppl.5): s.143-146 (endothelin-1); Hubbard et al. (1989), Annals Int. Med. 3: 206-12 (α1-antitrypsin); Smith et al. (1989), J. Clin. Invest. 84: 1145-6 (α1-proteinase); Oswein et al. (March 1990), "Aerosolization of Proteins", Proc. Symp. Resp. Drug Delivery II, Keystone, Colorado (recombinant human growth hormone); Debs et al. (1988), J. Immunol. 140: 3482-8 (interferon-γ and tumor necrosis factor α) and Platz et al., U.S. Patent No. 5,284,656 (granulocyte colony stimulating factor).

Contemplated for use in the practice of this invention are a wide range of mechanical devices designed for pulmonary delivery of therapeutic products, including but not limited to nebulizers, metered dose inhalers, and powder inhalers, all of which are familiar to those skilled in the art. Some specific examples of commercially available devices suitable for the practice of this invention are the Ultravent nebulizer, manufactured by Mallinckrodt, Inc., St. Louis, Missouri; the Acorn II nebulizer, manufactured by Marquest Medical Products, Englewood, Colorado; the Ventolin metered dose inhaler, manufactured by Glaxo Inc., Research Triangle Park, North Carolina; and the Spinhaler powder inhaler, manufactured by Fisons Corp., Bedford, Massachusetts.

All such devices require the use of formulations suitable for the dispensing of the inventive compound. Typically, each formulation is specific to the type of device employed and may involve the use of an appropriate propellant material, in addition to diluents, adjuvants and/or carriers useful in therapy.

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The inventive compound should most advantageously be prepared in particulate form with an average particle size of less than 10 μm (or microns), most preferably 0.5 to 5 μm , for most effective delivery to the distal lung.

Pharmaceutically acceptable carriers include carbohydrates such as trehalose, mannitol, xylitol, sucrose, lactose, and sorbitol. Other ingredients for use in formulations may include DPPC, DOPE, DSPC and DOPC. Natural or synthetic surfactants may be used. PEG may be used (even apart from its use in derivatizing the protein or analog). Dextrans, such as cyclodextran, may be used. Bile salts and other related enhancers may be used. Cellulose and cellulose derivatives may be used. Amino acids may be used, such as use in a buffer formulation.

Also, the use of liposomes, microcapsules or microspheres, inclusion complexes, or other types of carriers is contemplated.

Formulations suitable for use with a nebulizer, either jet or ultrasonic, will typically comprise the inventive compound dissolved in water at a concentration of about 0.1 to 25 mg of biologically active protein per mL of solution. The formulation may also include a buffer and a simple sugar (e.g., for protein stabilization and regulation of osmotic pressure). The nebulizer formulation may also contain a surfactant, to reduce or prevent surface induced aggregation of the protein caused by atomization of the solution in forming the aerosol.

Formulations for use with a metered-dose inhaler device will generally comprise a finely divided powder containing the inventive

compound suspended in a propellant with the aid of a surfactant. The propellant may be any conventional material employed for this purpose, such as a chlorofluorocarbon, a hydrochlorofluorocarbon, a hydrocarbon, including trichlorofluoromethane, dichlorodifluoromethane, dichlorotetrafluoroethanol, and 1,1,1,2-tetrafluoroethane, or combinations thereof. Suitable surfactants include sorbitan trioleate and soya lecithin. Oleic acid may also be useful as a surfactant.

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Formulations for dispensing from a powder inhaler device will comprise a finely divided dry powder containing the inventive compound and may also include a bulking agent, such as lactose, sorbitol, sucrose, mannitol, trehalose, or xylitol in amounts which facilitate dispersal of the powder from the device, e.g., 50 to 90% by weight of the formulation.

Nasal delivery forms. Nasal delivery of the inventive compound is also contemplated. Nasal delivery allows the passage of the protein to the blood stream directly after administering the therapeutic product to the nose, without the necessity for deposition of the product in the lung. Formulations for nasal delivery include those with dextran or cyclodextran. Delivery via transport across other mucous membranes is also contemplated.

<u>Dosages</u>. The dosage regimen involved in a method for treating the above-described conditions will be determined by the attending physician, considering various factors which modify the action of drugs, e.g. the age, condition, body weight, sex and diet of the patient, the severity of any infection, time of administration and other clinical factors. Generally, the daily regimen should be in the range of 0.1-1000 micrograms of the inventive compound per kilogram of body weight, preferably 0.1-150 micrograms per kilogram.

Specific preferred embodiments

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The inventors have determined preferred peptide sequences for molecules having many different kinds of activity. The inventors have further determined preferred structures of these preferred peptides combined with preferred linkers and vehicles. Preferred structures for these preferred peptides listed in Table 21 below.

Table 21—Preferred embodiments

Sequence/structure	SEQ	Activity
•	ID	
: :	NO:	•
F1-(G),-IEGPTLRQWLAARA-(G),-IEGPTLRQWLAARA	337	TPO-mimetic
IEGPTLRQWLAARA-(G),-IEGPTLRQWLAARA-(G),- F1	338	TPO-mimetic
F1-(G)5-IEGPTLRQWLAARA		TPO-mimetic
	1032	
IEGPTLRQWLAARA -(G)₅- F¹		TPO-mimetic
	1033	
F'-(G),-GGTYSCHFGPLTWVCKPQGG-(G),-	339	EPO-mimetic
GGTYSCHFGPLTWVCKPQGG	ļl	
GGTYSCHFGPLTWVCKPQGG-(G),-		EPO-mimetic
GGTYSCHFGPLTWVCKPQGG-(G) _s -F'	340	EDO minorio
GGTYSCHFGPLTWVCKPQGG-(G) ₅ -F'		EPO-mimetic
	1034	71.15
F1-(G),-DFLPHYKNTSLGHRP	1045	TNF-α inhibitor
	1043	TNF-α inhibitor
DFLPHYKNTSLGHRP-(G) _s -F'	1046	ו ואר-מ ווווווטונטו
	1040	IL-1 R antagonist
F¹-(G) ₅ - FEWTPGYWQPYALPL	1047	IL-1 IT amayonist
ESMEROVANORYALDI (C) E	10-17	IL-1 R antagonist
FEWTPGYWQPYALPL-(G) ₅ -F'	1048	iz i i i anagomor
F¹-(G) ₅ -VEPNCDIHVMWEWECFERL	10.0	VEGF-antagonist
F-(G) ₅ -VEPNODIHVIVIVVEVVEOFERE	1049	, 120, C. M. 29,
VEPNCDIHVMWEWECFERL-(G) ₅ -F'	 	VEGF-antagonist
AELIAODII IAIMAAEAAFOI FILIF-/O/2-1	1050	
F'-(G) _s -CTTHWGFTLC	+	MMP inhibitor
in -(a) ₆ -of filwar teo	1051	
CTTHWGFTLC-(G) ₅ -F ¹		MMP inhibitor
011111101111011111111111111111111111111	1052	

[&]quot;F1" is an Fc domain as defined previously herein.

"Working examples

The compounds described above may be prepared as described below. These examples comprise preferred embodiments of the invention and are illustrative rather than limiting.

Example 1

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TPO-Mimetics

The following example uses peptides identified by the numbers appearing in Table A hereinafter.

Preparation of peptide 19. Peptide 17b (12 mg) and MeO-PEG-SH 5000 (30 mg, 2 equiv.) were dissolved in 1 ml aqueous buffer (pH 8). The mixture was incubated at RT for about 30 minutes and the reaction was checked by analytical HPLC, which showed a > 80% completion of the reaction. The pegylated material was isolated by preparative HPLC.

Preparation of peptide 20. Peptide 18 (14 mg) and MeO-PEG-maleimide (25 mg) were dissolved in about 1.5 ml aqueous buffer (pH 8). The mixture was incubated at RT for about 30 minutes, at which time about 70% transformation was complete as monitored with analytical HPLC by applying an aliquot of sample to the HPLC column. The pegylated material was purified by preparative HPLC.

Bioactivity assay. The TPO in vitro bioassay is a mitogenic assay utilizing an IL-3 dependent clone of murine 32D cells that have been transfected with human mpl receptor. This assay is described in greater detail in WO 95/26746. Cells are maintained in MEM medium containing 10% Fetal Clone II and 1 ng/ml mIL-3. Prior to sample addition, cells are prepared by rinsing twice with growth medium lacking mIL-3. An extended twelve point TPO standard curve is prepared, ranging from 33 to 39 pg/ml. Four dilutions, estimated to fall within the linear portion of the standard curve, (100 to 125 pg/ml), are prepared for each sample and run in triplicate. A volume of 100 µl of each dilution of sample or standard is added to appropriate wells of a 96 well microtiter plate

containing 10,000 cells/well. After forty-four hours at 37 °C and 10% CO₂, MTS (a tetrazolium compound which is bioreduced by cells to a formazan) is added to each well. Approximately six hours later, the optical density is read on a plate reader at 490 nm. A dose response curve (log TPO concentration vs. O.D.- Background) is generated and linear regression analysis of points which fall in the linear portion of the standard curve is performed. Concentrations of unknown test samples are determined using the resulting linear equation and a correction for the dilution factor.

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TMP tandem repeats with polyglycine linkers. Our design of sequentially linked TMP repeats was based on the assumption that a dimeric form of TMP was required for its effective interaction with c-Mpl (the TPO receptor) and that depending on how they were wound up against each other in the receptor context, the two TMP molecules could be tethered together in the C- to N-terminus configuration in a way that would not perturb the global dimeric conformation. Clearly, the success of the design of tandem linked repeats depends on proper selection of the length and composition of the linker that joins the C- and N-termini of the two sequentially aligned TMP monomers. Since no structural information of the TMP bound to c-Mpl was available, a series of repeated peptides with linkers composed of 0 to 10 and 14 glycine residues (Table A) were synthesized. Glycine was chosen because of its simplicity and flexibility, based on the rationale that a flexible polyglycine peptide chain might allow for the free folding of the two tethered TMP repeats into the required conformation, while other amino acid sequences may adopt undesired secondary structures whose rigidity might disrupt the correct packing of the repeated peptide in the receptor context.

The resulting peptides are readily accessible by conventional solid phase peptide synthesis methods (Merrifield (1963), <u>J. Amer. Chem. Soc.</u> 85: 2149) with either Fmoc or t-Boc chemistry. Unlike the synthesis of the

C-terminally linked parallel dimer which required the use of an orthogonally protected lysine residue as the initial branch point to build the two peptide chains in a pseudosymmetrical way (Cwirla et al. (1997), Science 276: 1696-9), the synthesis of these tandem repeats was a straightforward, stepwise assembly of the continuous peptide chains from the C- to N-terminus. Since dimerization of TMP had a more dramatic effect on the proliferative activity than binding affinity as shown for the Cterminal dimer (Cwirla et al. (1997)), the synthetic peptides were tested directly for biological activity in a TPO-dependent cell-proliferation assay using an IL-3 dependent clone of murine 32D cells transfected with the full-length c-Mpl (Palacios et al.,. Cell 41:727 (1985)). As the test results showed, all the polyglycine linked tandem repeats demonstrated >1000 fold increases in potency as compared to the monomer, and were even more potent than the C-terminal dimer in this cell proliferation assay. The absolute activity of the C-terminal dimer in our assay was lower than that of the native TPO protein, which is different from the previously reported findings in which the C-terminal dimer was found to be as active as the natural ligand (Cwirla et al. (1997)). This might be due to differences in the conditions used in the two assays. Nevertheless, the difference in activity between tandem (C terminal of first monomer linked to N terminal of second monomer) and C-terminal (C terminal of first monomer linked to C terminal of second monomer; also referred to as parallel) dimers in the same assay clearly demonstrated the superiority of tandem repeat strategy over parallel peptide dimerization. It is interesting to note that a wide range of length is tolerated by the linker. The optimal linker between tandem peptides with the selected TMP monomers apparently is composed of 8 glycines.

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Other tandem repeats. Subsequent to this first series of TMP tandem repeats, several other molecules were designed either with

different linkers or containing modifications within the monomer itself. The first of these molecules, peptide 13, has a linker composed of GPNG, a sequence known to have a high propensity to form a β -turn-type secondary structure. Although still about 100-fold more potent than the monomer, this peptide was found to be >10-fold less active than the equivalent GGGG-linked analog. Thus, introduction of a relatively rigid β -turn at the linker region seemed to have caused a slight distortion of the optimal agonist conformation in this short linker form.

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The Trp9 in the TMP sequence is a highly conserved residue among the active peptides isolated from random peptide libraries. There is also a 10 highly conserved Trp in the consensus sequences of EPO mimetic peptides and this Trp residue was found to be involved in the formation of a hydrophobic core between the two EMPs and contributed to hydrophobic interactions with the EPO receptor. Livnah et al. (1996), Science 273: 464-71). By analogy, the Trp9 residue in TMP might have a similar function in 15 dimerization of the peptide ligand, and as an attempt to modulate and estimate the effects of noncovalent hydrophobic forces exerted by the two indole rings, several analogs were made resulting from mutations at the Trp. So in peptide 14, the Trp residue was replaced in each of the two TMP monomers with a Cys, and an intramolecular disulfide bond was 20 formed between the two cysteines by oxidation which was envisioned to mimic the hydrophobic interactions between the two Trp residues in peptide dimerization. Peptide 15 is the reduced form of peptide 14. In peptide 16, the two Trp residues were replaced by Ala. As the assay data show, all three analogs were inactive. These data further demonstrated 25 that Trp is critical for the activity of the TPO mimetic peptide, not just for dimer formation.

The next two peptides (peptide 17a, and 18) each contain in their 8-amino acid linker a Lys or Cys residue. These two compounds are

precursors to the two PEGylated peptides (peptide 19 and 20) in which the side chain of the Lys or Cys is modified by a PEG moiety. A PEG moiety was introduced at the middle of a relatively long linker, so that the large PEG component (5 kDa) is far enough away from the critical binding sites in the peptide molecule. PEG is a known biocompatible polymer which is increasingly used as a covalent modifier to improve the pharmacokinetic profiles of peptide- and protein-based therapeutics.

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A modular, solution-based method was devised for convenient PEGylation of synthetic or recombinant peptides. The method is based on the now well established chemoselective ligation strategy which utilizes the specific reaction between a pair of mutually reactive functionalities. So, for pegylated peptide 19, the lysine side chain was preactivated with a bromoacetyl group to give peptide 17b to accommodate reaction with a thiol-derivatized PEG. To do that, an orthogonal protecting group, Dde, was employed for the protection of the lysine ε-amine. Once the whole peptide chain was assembled, the N-terminal amine was reprotected with t-Boc. Dde was then removed to allow for the bromoacetylation. This strategy gave a high quality crude peptide which was easily purified using conventional reverse phase HPLC. Ligation of the peptide with the thiolmodified PEG took place in aqueous buffer at pH 8 and the reaction completed within 30 minutes. MALDI-MS analysis of the purified, pegylated material revealed a characteristic, bell-shaped spectrum with an increment of 44 Da between the adjacent peaks. For PEG-peptide 20, a cysteine residue was placed in the linker region and its side chain thiol group would serve as an attachment site for a maleimide-containing PEG. Similar conditions were used for the pegylation of this peptide. As the assay data revealed, these two pegylated peptides had even higher in vitro bioactivity as compared to their unpegylated counterparts.

Peptide 21 has in its 8-amino acid linker a potential glycosylation motif, NGS. Since our exemplary tandem repeats are made up of natural amino acids linked by peptide bonds, expression of such a molecule in an appropriate eukaryotic cell system should produce a glycopeptide with the carbohydrate moiety added on the side chain carboxyamide of Asn. Glycosylation is a common post-translational modification process which can have many positive impacts on the biological activity of a given protein by increasing its aqueous solubility and in vivo stability. As the assay data show, incorporation of this glycosylation motif into the linker maintained high bioactivity. The synthetic precursor of the potential glycopeptide had in effect an activity comparable to that of the -(G)₈-linked analog. Once glycosylated, this peptide is expected to have the same order of activity as the pegylated peptides, because of the similar chemophysical properties exhibited by a PEG and a carbohydrate moiety.

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The last peptide is a dimer of a tandem repeat. It was prepared by oxidizing peptide 18, which formed an intermolecular disulfide bond between the two cysteine residues located at the linker. This peptide was designed to address the possibility that TMP was active as a tetramer. The assay data showed that this peptide was not more active than an average tandem repeat on an adjusted molar basis, which indirectly supports the idea that the active form of TMP is indeed a dimer, otherwise dimerization of a tandem repeat would have a further impact on the bioactivity.

In order to confirm the in vitro data in animals, one pegylated TMP tandem repeat (compound 20 in Table A) was delivered subcutaneously to normal mice via osmotic pumps. Time and dose-dependent increases were seen in platelet numbers for the duration of treatment. Peak platelet levels over 4-fold baseline were seen on day 8. A dose of $10^{\circ}\mu g/kg/day$ of the pegylated TMP repeat produced a similar response to rHuMGDF (non-pegylated) at $100 \, \mu g/kg/day$ delivered by the same route.

Table A—TPO-mimetic Peptides

Peptide	Compound	SEQ ID	Relative				
No.		NO:	Potency				
	TPO		++++				
	TMP monomer	13	+				
	TMP C-C dimer		+++-				
TMP-(G) _n -	TMP:						
1	n = 0	341	++++-				
2	n = 1	342	++++				
3	n = 2	343	++++				
4	n = 3	344	++++				
5	n = 4	345	++++				
6	n = 5	346	++++				
7	n = 6	347	++++				
8	n = 7	348	++++				
9	n = 8	349	++++-				
10	n = 9	350	++++				
11	n = 10	351	++++				
12	n = 14 _.	352	++++				
13	TMP-GPNG-TMP	353	+++				
14	IEGPTLRQCLAARA-GGGGGGGG-IEGPTLRQCLAARA	354					
15	(cyclic) IEGPTLRQCLAARA-GGGGGGGG- IEGPTLRQCLAARA (linear)	355	•				
16		356	_				
10	IEGPTLRQ <u>A</u> LAARA-GGGGGGGG- 356 - IEGPTLRQ <u>A</u> LAARA						
17a	TMP-GGGKGGGG-TMP	357	++++				
17a 17b	TMP-GGGK(BrAc)GGGG-TMP 358 N						
175	TMP-GGGCGGGG-TMP 359 ++						
19	TMP-GGGK(PEG)GGGG-TMP 360 ++++						
20	TMP-GGGC(PEG)GGGG-TMP 361 ++++						
21	TMP-GGGN*GSGG-TMP 362 ++++						
22	TMP-GGGCGGGG-TMP	363-	- ,				
	 TMP-GGCGGGG-TMP	363	++++				

<u>Discussion</u>. It is well accepted that MGDF acts in a way similar to hGH, i.e., one molecule of the protein ligand binds two molecules of the receptor for its activation. Wells <u>et al.</u>(1996), <u>Ann. Rev. Biochem.</u> 65: 609-34. Now, this interaction is mimicked by the action of a much smaller peptide, TMP. However, the present studies suggest that this mimicry requires the concerted action of two TMP molecules, as covalent dimerization of TMP in either a C-C parallel or C-N sequential fashion increased the <u>in vitro</u> biological potency of the original monomer by a factor of greater than 10³. The relatively low biopotency of the monomer is probably due to inefficient formation of the noncovalent dimer. A preformed covalent repeat has the ability to eliminate the entropy barrier for the formation of a noncovalent dimer which is exclusively driven by weak, noncovalent interactions between two molecules of the small, 14-residue peptide.

It is intriguing that this tandem repeat approach had a similar effect on enhancing bioactivity as the reported C-C dimerization is intriguing. These two strategies brought about two very different molecular configurations. The C-C dimer is a quasi-symmetrical molecule, while the tandem repeats have no such symmetry in their linear structures. Despite this difference in their primary structures, these two types of molecules appeared able to fold effectively into a similar biologically active conformation and cause the dimerization and activation of c-Mpl. These experimental observations provide a number of insights into how the two TMP molecules may interact with one another in binding to c-Mpl. First, the two C-termini of the two bound TMP molecules must be in relatively close proximity with each other, as suggested by data on the C-terminal dimer. Second, the respective N- and C-termini of the two TMP molecules in the receptor complex must also be very closely aligned with each other, such that they can be directly tethered together with a single peptide bond

to realize the near maximum activity-enhancing effect brought about by the tandem repeat strategy. Insertion of one or more (up to 14) glycine residues at the junction did not increase (or decrease) significantly the activity any further. This may be due to the fact that a flexible polyglycine peptide chain is able to loop out easily from the junction without causing any significant changes in the overall conformation. This flexibility seems to provide the freedom of orientation for the TMP peptide chains to fold into the required conformation in interacting with the receptor and validate it as a site of modification. Indirect evidence supporting this came from the study on peptide 13, in which a much more rigid b-turnforming sequence as the linker apparently forced a deviation of the backbone alignment around the linker which might have resulted in a slight distortion of the optimal conformation, thus resulting in a moderate (10-fold) decrease in activity as compared with the analogous compound with a 4-Gly linker. Third, Trp9 in TMP plays a similar role as Trp13 in EMP, which is involved not only in peptide:peptide interaction for the formation of dimers but also is important for contributing hydrophobic forces in peptide:receptor interaction. Results obtained with the W to C mutant analog, peptide 14, suggest that a covalent disulfide linkage is not sufficient to approximate the hydrophobic interactions provided by the Trp pair and that, being a short linkage, it might bring the two TMP monomers too close, therefore perturbing the overall conformation of the optimal dimeric structure.

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An analysis of the possible secondary structure of the TMP peptide can provide further understanding on the interaction between TMP and c-Mpl. This can be facilitated by making reference to the reported structure of the EPO mimetic peptide. Livnah et al. (1996), Science 273:464-75 The receptor-bound EMP has a b-hairpin structure with a b-turn formed by the highly consensus Gly-Pro-Leu-Thr at the center of its sequence. Instead of

GPLT, TMP has a highly selected GPTL sequence which is likely to form a similar turn. However, this turn-like motif is located near the N-terminal part in TMP. Secondary structure prediction using Chau-Fasman method suggests that the C-terminal half of the peptide has a tendency to adopt a helical conformation. Together with the highly conserved Trp at position 9, this C-terminal helix may contribute to the stabilization of the dimeric structure. It is interesting to note that most of our tandem repeats are more potent than the C-terminal parallel dimer. Tandem repeats seem to give the molecule a better fit conformation than does the C-C parallel dimerization. The seemingly asymmetric feature of a tandem repeat might have brought it closer to the natural ligand which, as an asymmetric molecule, uses two different sites to bind two identical receptor molecules.

Introduction of a PEG moiety was envisaged to enhance the <u>in vivo</u> activity of the modified peptide by providing it a protection against proteolytic degradation and by slowing down its clearance through renal filtration. It was unexpected that pegylation could further increase the <u>in vitro</u> bioactivity of a tandem repeated TMP peptide in the cell-based proliferation assay.

Example 2

Fc-TMP fusions

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TMPs (and EMPs as described in Example 3) were expressed in either monomeric or dimeric form as either N-terminal or C-terminal fusions to the Fc region of human IgG1. In all cases, the expression construct utilized the luxPR promoter promoter in the plasmid expression vector pAMG21.

Fc-TMP. A DNA sequence coding for the Fc region of human IgG1 fused in-frame to a monomer of the TPO-mimetic peptide was constructed using standard PCR technology. Templates for PCR reactions were the pFc-A3 vector and a synthetic TMP gene. The synthetic gene was

constructed from the 3 overlapping oligonucleotides (SEQ ID NOS: 364, 365, and 366, respectively) shown below:

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1842-97 AAA AAA GGA TCC TCG AGA TTA AGC ACG AGC AGC CAG CCA

5 1842-98 AAA GGT GGA GGT GGT ATC GAA GGT CCG ACT CTG CGT
1842-99 CAG TGG CTG GCT GCT CGT GCT TAA TCT CGA GGA TCC TTT
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These oligonucleotides were annealed to form the duplex encoding an amino acid sequence (SEQ ID NOS: 367 and 368, respectively) shown below:

This duplex was amplified in a PCR reaction using 1842-98 and 1842-97 as the sense and antisense primers.

The Fc portion of the molecule was generated in a PCR reaction with pFc-A3 using the primers shown below (SEQ ID NOS: 369 and 370):

1216-52

AAC ATA AGT ACC TGT AGG ATC G

1830-51

TTCGATACCA CCACCTCCAC CTTTACCCGG AGACAGGGAG AGGCTCTTCTGC

The oligonucleotides 1830-51 and 1842-98 contain an overlap of 24

nucleotides, allowing the two genes to be fused together in the correct reading frame by combining the above PCR products in a third reaction using the outside primers, 1216-52 and 1842-97.

The final PCR gene product (the full length fusion gene) was digested with restriction endonucleases XbaI and BamHI, and then ligated into the vector pAMG21 and transformed into competent <u>E. coli</u> strain 2596 cells as described for EMP-Fc herein. Clones were screened for the ability to produce the recombinant protein product and to possess the

gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #3728.

The nucleotide and amino acid sequences (SEQ ID NOS: 5 and 6) of the fusion protein are shown in Figure 7.

Fc-TMP-TMP. A DNA sequence coding for the Fc region of human IgG1 fused in-frame to a dimer of the TPO-mimetic peptide was constructed using standard PCR technology. Templates for PCR reactions were the pFc-A3 vector and a synthetic TMP-TMP gene. The synthetic gene was constructed from the 4 overlapping oligonucleotides (SEQ ID

NOS: 371 to 374, respectively) shown below:

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The 4 oligonucleotides were annealed to form the duplex encoding an amino acid sequence (SEQ ID NOS: 375 and 376, respectively) shown below:

This duplex was amplified in a PCR reaction using 1830-52 and 1830-55 as the sense and antisense primers.

The Fc portion of the molecule was generated in a PCR reaction with pFc-A3 using the primers 1216-52 and 1830-51 as described above for

Fc-TMP. The full length fusion gene was obtained from a third PCR reaction using the outside primers 1216-52 and 1830-55.

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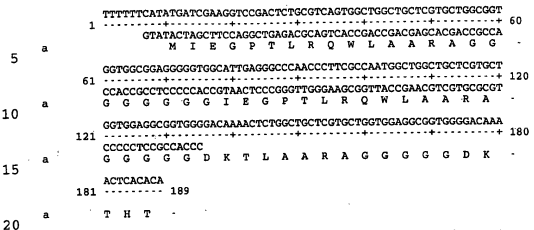
The final PCR gene product (the full length fusion gene) was digested with restriction endonucleases <u>XbaI</u> and <u>BamHI</u>, and then ligated into the vector pAMG21 and transformed into competent <u>E. coli</u> strain 2596 cells as described in example 1. Clones were screened for the ability to produce the recombinant protein product and to possess the gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #3727.

The nucleotide and amino acid sequences (SEQ ID NOS: 7 and 8) of the fusion protein are shown in Figure 8.

TMP-TMP-Fc. A DNA sequence coding for a tandem repeat of the TPO-mimetic peptide fused in-frame to the Fc region of human IgG1 was constructed using standard PCR technology. Templates for PCR reactions were the EMP-Fc plasmid from strain #3688 (see Example 3) and a synthetic gene encoding the TMP dimer. The synthetic gene for the tandem repeat was constructed from the 7 overlapping oligonucleotides shown below (SEQ ID NOS: 377 to 383, respectively):

20	1885-52	TTT 7	TTT	CAT	ATG	ATC	GAA	GGT	CCG	ACT	CTG	CGT	CAG	TGG
	1885-53	AGC A			AGC	CAG	CCA	CTG	ACG	CAG	AGT	CGG	ACC	TTC
25	1885-54	CTG C		GCT	CGT	GCT	GGT	GGA	GGC	GGT	GGG	GAC	AAA	ACT
2.0	1885-55	CTG C				GCT	GGC	GGT	GGT	GGC	GGA	GGG	GGT	GGC
30	1885-56	AAG (GGT	TGG	GCC	CTC	AAT	GCC	ACC	CCC
35	1885-57	ACC (CTT	GCA	GCA	CGC	GCA	GGG	GGA	GGC
	1885-58	CCC 2	ACC	GCC	TCC	CCC	TGC	GCG	TGC	TGC				

These oligonucleotides were annealed to form the duplex shown encoding an amino acid sequence shown below (SEQ ID NOS 384 and 385):



This duplex was amplified in a PCR reaction using 1885-52 and 1885-58 as the sense and antisense primers.

The Fc portion of the molecule was generated in a PCR reaction with DNA from the EMP-Fc fusion strain #3688 (see Example 3) using the primers 1885-54 and 1200-54. The full length fusion gene was obtained from a third PCR reaction using the outside primers 1885-52 and 1200-54.

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The final PCR gene product (the full length fusion gene) was digested with restriction endonucleases XbaI and BamHI, and then ligated into the vector pAMG21 and transformed into competent E. coli strain 2596 cells as described for Fc-EMP herein. Clones were screened for the ability to produce the recombinant protein product and to possess the gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #3798.

The nucelotide and amino acid sequences (SEQ ID NOS: 9 and 10)

of the fusion protein are shown in Figure 9.

TMP-Fc. A DNA sequence coding for a monomer of the TPO-mimetic peptide fused in-frame to the Fc region of human IgG1 was obtained fortuitously in the ligation in TMP-TMP-Fc, presumably due to the ability of primer 1885-54 to anneal to 1885-53 as well as to 1885-58. A single clone having the correct nucleotide sequence for the TMP-Fc construct was selected and designated Amgen strain #3788.

The nucleotide and amino acid sequences (SEQ ID NOS: 11 and 12) of the fusion protein are shown in Figure 10.

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Expression in E. coli. Cultures of each of the pAMG21-Fc-fusion constructs in E. coli GM221 were grown at 37 °C in Luria Broth medium containing 50 mg/ml kanamycin. Induction of gene product expression from the luxPR promoter was achieved following the addition of the synthetic autoinducer N-(3-oxohexanoyl)-DL-homoserine lactone to the culture media to a final concentration of 20 ng/ml. Cultures were incubated at 37 °C for a further 3 hours. After 3 hours, the bacterial cultures were examined by microscopy for the presence of inclusion bodies and were then collected by centrifugation. Refractile inclusion bodies were observed in induced cultures indicating that the Fc-fusions were most likely produced in the insoluble fraction in E. coli. Cell pellets were lysed directly by resuspension in Laemmli sample buffer containing 10% b-mercaptoethanol and were analyzed by SDS-PAGE. In each case, an intense coomassie-stained band of the appropriate molecular weight was observed on an SDS-PAGE gel.

pAMG21. The expression plasmid pAMG21 can be derived from the Amgen expression vector pCFM1656 (ATCC #69576) which in turn be derived from the Amgen expression vector system described in US Patent No. 4,710,473. The pCFM1656 plasmid can be derived from the described pCFM836 plasmid (Patent No. 4,710,473) by:

- (a) destroying the two endogenous <u>NdeI</u> restriction sites by end filling with T4 polymerase enzyme followed by blunt end ligation;
- (b) replacing the DNA sequence between the unique <u>AatII</u> and <u>ClaI</u> restriction sites containing the synthetic P_L promoter with a similar fragment obtained from pCFM636 (patent No. 4,710,473) containing the PL promoter (see SEQ ID NO: 386 below); and

(c) substituting the small DNA sequence between the unique <u>ClaI</u> and <u>KpnI</u> restriction sites with the oligonucleotide having the sequence of SEQ ID NO: 388.

SEQ ID NO: 386:

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- - 5 CGATTTGATTCTAGAAGGAGGAATAACATATGGTTAACGCGTTGGAATTCGGTAC
 3 TAAACTAAGATCTTCCTCCTTATTGTATACCAATTGCGCAACCTTAAGC
 5 Clai

 Kpnl
 - The expression plasmid pAMG21 can then be derived from pCFM1656 by making a series of site-directed base changes by PCR overlapping oligo mutagenesis and DNA sequence substitutions. Starting with the <u>BglII</u> site (plasmid bp # 180) immediately 5' to the plasmid replication promoter P_{copB} and proceeding toward the plasmid replication genes, the base pair changes are as shown in Table B below.

Table B—Base pair changes resulting in pAMG21

	pAMG21 bp #	bp in pCFM1656	bp changed to in pAMG21
5	# 204	T/A	C/G
	# 428	A/T	G/C
	# 509	G/C	A/T
	# 617	••	insert two G/C bp
	# 679	G/C	T/A
10	# 980	T/A	C/G
	# 994	G/C	A/ T
	# 1004	A/T	C/G
	# 1007	C/G	T/A
	# 1028	A/T	T/A
15	# 1047	C/G	T/A
	# 1178	G/C	T/A
	# 1466	G/C	T/A
	# 2028	G/C	bp deletion
	# 2187	C/G	T/A
20	# 2480	A/T	T/A
	# 2499-2502	AGTG	<u>GTCA</u>
		TCAC	CAGT
25	# 2642	TCCGAGC AGGCTCG	7 bp deletion
	# 3435	G/C	A/T
	# 3446	G/C	A/T .
30	# 3643	A/T	T/A
30		A/T	T/A

The DNA sequence between the unique <u>Aat</u>II (position #4364 in pCFM1656) and <u>Sac</u>II (position #4585 in pCFM1656) restriction sites is substituted with the DNA sequence (SEQ ID NO: 23) shown in Figures 17A and 17B. During the ligation of the sticky ends of this substitution DNA sequence, the outside <u>Aat</u>II and <u>Sac</u>II sites are destroyed. There are unique AatII and <u>Sac</u>II sites in the substituted DNA.

GM221 (Amgen #2596). The Amgen host strain #2596 is an E.coli K-12 strain derived from Amgen strain #393. It has been modified to contain both the temperature sensitive lambda repressor cI857s7 in the early ebg region and the lacl^Q repressor in the late ebg region (68 minutes). The presence of these two repressor genes allows the use of this host with a variety of expression systems, however both of these repressors are irrelevant to the expression from luxP_R. The untransformed host has no antibiotic resistances.

The ribosome binding site of the cI857s7 gene has been modified to include an enhanced RBS. It has been inserted into the <u>ebg</u> operon between nucleotide position 1170 and 1411 as numbered in Genbank accession number M64441Gb_Ba with deletion of the intervening <u>ebg</u> sequence. The sequence of the insert is shown below with lower case letters representing the <u>ebg</u> sequences flanking the insert shown below (SEQ ID NO: 388):

The construct was delivered to the chromosome using a recombinant phage called MMebg-cI857s7enhanced RBS #4 into F'tet/393. After recombination and resolution only the chromosomal insert described

above remains in the cell. It was renamed F'tet/GM101. F'tet/GM101 was then modified by the delivery of a lacI^Q construct into the <u>ebg</u> operon between nucleotide position 2493 and 2937 as numbered in the Genbank accession number M64441Gb_Ba with the deletion of the intervening <u>ebg</u> sequence. The sequence of the insert is shown below with the lower case letters representing the <u>ebg</u> sequences flanking the insert (SEQ ID NO: 389) shown below:

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ggcggaaaccGACGTCCATCGAATGGTGCAAAACCTTTCGCGGTATGGCATGATAGCGCCCGGAAGAGAGTCA ATTCAGGGTGGTGAATGTGAAACCAGTAACGTTATACGATGTCGCAGAGTATGCCGGTGTCTCTTATCAGACC 10 GTTTCCCGCGTGGTGAACCAGGCCAGCCACGTTTCTGCGAAAACGCGGGAAAAAGTCGAAGCGGCGATGGCGG AGCTGAATTACATTCCCAACCGCGTGGCACAACAACTGGCGGCAAACAGTCGCTCCTGATTGGCGTTGCCAC CTCCAGTCTGGCCCTGCACGCGCCGTCGCAAATTGTCGCGGCGATTAAATCTCGCGCCGATCAACTGGGTGCC AGCGTGGTGGTGTCGATGGTAGAACGAAGCGGCGTCGAAGCCTGTAAAGCGGCGGTGCACAATCTTCTCGCGC TAATGTTCCGGCGTTATTTCTTGATGTCTCTGACCAGACACCCATCAACAGTATTATTTCTCCCATGAAGAC 15 GGTACGCGACTGGGCGTGGAGCATCTGGTCGCATTGGGTCACCAGCAAATCGCGCTGTTAGCGGGCCCCATTAA GTTCTGTCTCGGCGCGTCTGCGTCTGGCTGGCTGGCATAAATATCTCACTCGCAATCAAATTCAGCCGATAGC GGAACGGGAAGGCGACTGGAGTGCCATGTCCGGTTTTCAACAAACCATGCAAATGCTGAATGAGGGCATCGTT CCCACTGCGATGCTGGTTGCCAACGATCAGATGGCGCTGGGCGCAATGCGCGCCATTACCGAGTCCGGGCTGC GCGTTGGTGCGGATATCTCGGTAGTGGGATACGACGATACCGAAGACAGCTCATGTTATATCCCGCCGTTAAC 20 CACCATCAAACAGGATTTTCGCCTGCTGGGGCAAACCAGCGTGGACCGCTTGCTGCAACTCTCTCAGGGCCAG GCGGTGAAGGGCAATCAGCTGTTGCCCGTCTCACTGGTGAAAAGAAAAACCACCCTGGCGCCCCAATACGCAAA CCGCCTCTCCCCGCGCGTTGGCCGATTCATTAATGCAGCTGGCACGACAGGTTTCCCGACTGGAAAGCGGACA GTAAGGTACCATAGGATCCaggcacagga 25

The construct was delivered to the chromosome using a recombinant phage called AGebg-LacIQ#5 into F'tet/GM101. After recombination and resolution only the chromosomal insert described above remains in the cell. It was renamed F'tet/GM221. The F'tet episome was cured from the strain using acridine orange at a concentration of 25 μ g/ml in LB. The cured strain was identified as tetracyline sensitive and was stored as GM221.

Expression. Cultures of pAMG21-Fc-TMP-TMP in *E. coli* GM221 in

Luria Broth medium containing 50 μg/ml kanamycin were incubated at

37°C prior to induction. Induction of Fc-TMP-TMP gene product

expression from the luxPR promoter was achieved following the addition

of the synthetic autoinducer N-(3-oxohexanoyl)-DL-homoserine lactone to

the culture media to a final concentration of 20 ng/ml and cultures were

incubated at 37°C for a further 3 hours. After 3 hours, the bacterial

cultures were examined by microscopy for the presence of inclusion bodies and were then collected by centrifugation. Refractile inclusion bodies were observed in induced cultures indicating that the Fc-TMP-TMP was most likely produced in the insoluble fraction in *E. coli*. Cell pellets were lysed directly by resuspension in Laemmli sample buffer containing 10% •-mercaptoethanol and were analyzed by SDS-PAGE. An intense Coomassie stained band of approximately 30kDa was observed on an SDS-PAGE gel. The expected gene product would be 269 amino acids in length and have an expected molecular weight of about 29.5 kDa. Fermentation was also carried out under standard batch conditions at the

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Fermentation was also carried out under standard batch conditions at the 10 L scale, resulting in similar expression levels of the Fc-TMP-TMP to those obtained at bench scale.

Purification of Fc-TMP-TMP. Cells are broken in water (1/10) by high pressure homogenization (2 passes at 14,000 PSI) and inclusion bodies are harvested by centrifugation (4200 RPM in J-6B for 1 hour). Inclusion bodies are solubilized in 6M guanidine, 50mM Tris, 8mM DTT, pH 8.7 for 1 hour at a 1/10 ratio. The solubilized mixture is diluted 20 times into 2M urea, 50 mM tris, 160mM arginine, 3mM cysteine, pH 8.5. The mixture is stirred overnight in the cold and then concentrated about 10 fold by ultafiltration. It is then diluted 3 fold with 10mM Tris, 1.5M urea, pH 9. The pH of this mixture is then adjusted to pH 5 with acetic acid. The precipitate is removed by centrifugation and the supernatant is loaded onto a SP-Sepharose Fast Flow column equilibrated in 20mM NaAc, 100 mM NaCl, pH 5(10mg/ml protein load, room temperature). The protein is eluted off using a 20 column volume gradient in the same buffer ranging from 100mM NaCl to 500mM NaCl. The pool from the column is diluted 3 fold and loaded onto a SP-Sepharose HP column in 20 mM NaAc, 150 mM NaCl, pH 5(10 mg/ml protein load, room temperature). The protein is eluted off using a 20 column volume gradient

in the same buffer ranging from 150 mM NaCl to 400 mM NaCl. The peak is pooled and filtered.

<u>Characterization of Fc-TMP activity</u>. The following is a summary of <u>in vivo</u> data in mice with various compounds of this invention.

Mice: Normal female BDF1 approximately 10-12 weeks of age.

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Bleed schedule: Ten mice per group treated on day 0, two groups started 4 days apart for a total of 20 mice per group. Five mice bled at each time point, mice were bled a minimum of three times a week. Mice were anesthetized with isoflurane and a total volume of 140-160 µl of blood was obtained by puncture of the orbital sinus. Blood was counted on a Technicon H1E blood analyzer running software for murine blood. Parameters measured were white blood cells, red blood cells, hematocrit, hemoglobin, platelets, neutrophils.

treatment or implanted with 7-day micro-osmotic pumps for continuous delivery. Subcutaneous injections were delivered in a volume of 0.2 ml. Osmotic pumps were inserted into a subcutaneous incision made in the skin between the scapulae of anesthetized mice. Compounds were diluted in PBS with 0.1% BSA. All experiments included one control group,

labeled "carrier" that were treated with this diluent only. The concentration of the test articles in the pumps was adjusted so that the calibrated flow rate from the pumps gave the treatment levels indicated in the graphs.

Compounds: A dose titration of the compound was delivered to mice in 7 day micro-osmotic pumps. Mice were treated with various compounds at a single dose of 100 µg/kg in 7 day osmotic pumps. Some of the same compounds were then given to mice as a single bolus injection.

Activity test results: The results of the activity experiments are shown in Figures 11 and 12. In dose response assays using 7-day micro-

osmotic pumps, the maximum effect was seen with the compound of SEQ ID NO: 18 was at 100 μ g/kg/day; the 10 μ g/kg/day dose was about 50% maximally active and 1 μ g/kg/day was the lowest dose at which activity could be seen in this assay system. The compound at 10 μ g/kg/day dose was about equally active as 100 μ g/kg/day unpegylated rHu-MGDF in the same experiment.

Example 3

Fc-EMP fusions

Fc-EMP. A DNA sequence coding for the Fc region of human IgG1 fused in-frame to a monomer of the EPO-mimetic peptide was constructed using standard PCR technology. Templates for PCR reactions were a vector containing the Fc sequence (pFc-A3, described in International application WO 97/23614, published July 3, 1997) and a synthetic gene encoding EPO monomer. The synthetic gene for the monomer was constructed from the 4 overlapping oligonucleotides (SEQ ID NOS: 390 to

10 393, respectively) shown below:

```
1798-2 TAT GAA AGG TGG AGG TGG TGG AGG TAC TTA CTC TTG
CCA CTT CGG CCC GCT GAC TTG G

1798-3 CGG TTT GCA AAC CCA AGT CAG CGG GCC GAA GTG GCA AGA
TTG CTC TCC ACC TTC GCA GCT TCC ACC TTT CAT

1798-4 GTT TGC AAA CCG CAG GGT GGC GGC GGC GGC GGC GGT GGT
ACC TAT TCC TGT CAT TTT

1798-5 CCA GGT CAG CGG GCC AAA ATG ACA GGA ATA GGT ACC ACC
CTG CCC GCC GCC GCC GCC CTG
```

The 4 oligonucleotides were annealed to form the duplex encoding an amino acid sequence (SEQ ID NOS: 394 and 395, respectively) shown below:

This duplex was amplified in a PCR reaction using

```
40 1798-18 GCA GAA GAG CCT CTC CCT GTC TCC GGG TAA AGG TGG AGG TGG TGG AGG TAC TTA CTC TC TC GGG TAA AGG TGG TGG AGG TAC TTA AGG TGG TGG AGG TAC TTA
```

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1798-19
CTA ATT GGA TCC ACG AGA TTA ACC ACC
CTG CGG TTT GCA A

as the sense and antisense primers (SEQ ID NOS: 396 and 397, respectively).

The Fc portion of the molecule was generated in a PCR reaction with pFc-A3 using the primers

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1216-52
AAC ATA AGT ACC TGT AGG ATC G
1798-17
AGA GTA AGT ACC TCC ACC ACC ACC TCC ACC TTT ACC CGG
AGA CAG GGA GAG GCT CTT CTG C

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which are SEQ ID NOS: 398 and 399, respectively. The oligonucleotides 1798-17 and 1798-18 contain an overlap of 61 nucleotides, allowing the two genes to be fused together in the correct reading frame by combining the above PCR products in a third reaction using the outside primers, 1216-52 and 1798-19.

The final PCR gene product (the full length fusion gene) was digested with restriction endonucleases XbaI and BamHI, and then ligated into the vector pAMG21 (described below), also digested with XbaI and BamHI. Ligated DNA was transformed into competent host cells of E. coli strain 2596 (GM221, described herein). Clones were screened for the ability to produce the recombinant protein product and to possess the gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #3718.

The nucleotide and amino acid sequence of the resulting fusion protein (SEQ ID NOS: 15 and 16) are shown in Figure 13.

EMP-Fc. A DNA sequence coding for a monomer of the EPO-mimetic peptide fused in-frame to the Fc region of human IgG1 was constructed using standard PCR technology. Templates for PCR reactions were the pFC-A3a vector and a synthetic gene encoding EPO monomer.

The synthetic gene for the monomer was constructed from the 4 overlapping oligonucleotides 1798-4 and 1798-5 (above) and 1798-6 and 1798-7 (SEQ ID NOS: 400 and 401, respectively) shown below:

PCT/US99/25044 WO 00/24782

```
1798-6 GGC CCG CTG ACC TGG GTA TGT AAG CCA CAA GGG GGT GGG
            GGA GGC GGG GGG TAA TCT CGA G
     1798-7 GAT CCT CGA GAT TAC CCC CCG CCT CCC CCA CCC CCT TGT
 5
            GGC TTA CAT AC
     The 4 oligonucleotides were annealed to form the duplex encoding an
     amino acid sequence (SEQ ID NOS: 402 and 403, respectively) shown
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     below:
              GTTTGCAAACCGCAGGGTGGCGGCGGCGGCGGTGGTACCTATTCCTGTCATTTTGGC
                          GTCCCACCGCCGCCGCCGCCACCATGGATAAGGACAGTAAAACCG
15
              V C K P Q G G G G G G T Y S C H F G
              CCGCTGACCTGGGTATGTAAGCCACAAGGGGGTGGGGGGAGGCGGGGGGTAATCTCGAG
              GGCGACTGGACCCATACATTCGGTGTTCCCCCACCCCCTCCGCCCCCATTAGAGCTCCTAG
20
              PLTWVCKPQGGGGGG
            This duplex was amplified in a PCR reaction using
                  TTA TTT CAT ATG AAA GGT GGT AAC TAT TCC TGT CAT TTT
     1798-21
25
     and
     1798-22
                  TGG ACA TGT GTG AGT TTT GTC CCC CCC GCC TCC CCC ACC
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     as the sense and antisense primers (SEQ ID NOS: 404 and 405,
     respectively).
            The Fc portion of the molecule was generated in a PCR reaction
     with pFc-A3 using the primers
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     1798-23
                  AGG GGG TGG GGG AGG CGG GGG GGA CAA AAC TCA CAC ATG
     and
40
     1200-54
                  GTT ATT GCT CAG CGG TGG CA
      which are SEQ ID NOS: 406 and 407, respectively. The oligonucleotides
      1798-22 and 1798-23 contain an overlap of 43 nucleotides, allowing the two
      genes to be fused together in the correct reading frame by combining the
     above PCR products in a third reaction using the outside primers, 1787-21
```

The final PCR gene product (the full length fusion gene) was digested with restriction endonucleases XbaI and BamHI, and then ligated

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and 1200-54.

into the vector pAMG21 and transformed into competent <u>E. coli</u> strain 2596 cells as described above. Clones were screened for the ability to produce the recombinant protein product and to possess the gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #3688.

The nucleotide and amino acid sequences (SEQ ID NOS: 17 and 18) of the resulting fusion protein are shown in Figure 14.

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EMP-EMP-Fc. A DNA sequence coding for a dimer of the EPO-mimetic peptide fused in-frame to the Fc region of human IgG1 was constructed using standard PCR technology. Templates for PCR reactions were the EMP-Fc plasmid from strain #3688 above and a synthetic gene encoding the EPO dimer. The synthetic gene for the dimer was constructed from the 8 overlapping oligonucleotides (SEQ ID NOS:408 to 415, respectively) shown below:

15	1869-23	TTT TAG							GAT	TTG	AGT	TTT	AAC	TTT
20	1869-48	TAA AA	AAG	TTA	AAA	CTC	AAA	TCT	AGA	ATC	AAA	TCG	ATA	AAA
	1871-72		GGT TGC			TCT	TGC	CAC	TTC	GGC	CCG	CTG	ACT	TGG
25	1871-73				GCC CTC			GCA	AGA	GTA	AGT	ACC	TCC	CAT
20	1871-74	CAG CAT	GGT TTT	GGC GGC	GGC CCG	GGC CTG	GGC ACC	GGC TGG	GGT	GGT	ACC	TAT	TCC	TGT
30	1871-75							ACC CCA		GCC	GCC	GCC	GCC	GCC
35	1871-78				CCA ACA			GGT	GGG	GGA	GGC	GGG	GGG	GAC
	1871-79	AGT ACA	TTT TAC	GTC CCA	CCC GGT	CCC CAG	GCC CGG	TCC GCC	CCC	ACC	ccc	TTG	TGG	CTT

The 8 oligonucleotides were annealed to form the duplex encoding an amino acid sequence (SEQ ID NOS: 416 and 417, respectively) shown below:

TTTTTTATCGATTTGATTCTAGATTTGAGTTTTAACTTTTAGAAGGAGGAATAAAATATG

1 ----+ 60
AAAAAATAGCTAAACTAAACTCAAAATTGAAAATCTTCCTCCTTATTTTATAC
M

		61																				TGGC	
_		61					GAG	AAC	GGT	GAA	GCC	GGG	CGA	CTG	AAC	CCA	AAC	GTT	TGG			ACCG	
5	a		G	G	T	Y	S	С	Н	F	G	P	L	T	W	V	С	K	P	Q	G	G	-
		121																				TAAG	
10	a		CC G	GCC G	GCC G	GCC G	GCC G	ACC G			AAG S		AGT. H							CCA V		ATTC K	-
		181																TCC		28			
15	a		GG'	TGT: Q	TCC G	CCC. G	ACC G	CCC G	TCC G	GCC G	CCC G	CCT D	GTT K	TTG T	A H	т	С	P	-				

This duplex was amplified in a PCR reaction using 1869-23 and 1871-79 (shown above) as the sense and antisense primers.

The Fc portion of the molecule was generated in a PCR reaction with strain 3688 DNA using the primers 1798-23 and 1200-54 (shown above).

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The oligonucleotides 1871-79 and 1798-23 contain an overlap of 31 nucleotides, allowing the two genes to be fused together in the correct reading frame by combining the above PCR products in a third reaction using the outside primers, 1869-23 and 1200-54.

The final PCR gene product (the full length fusion gene) was digested with restriction endonucleases XbaI and BamHI, and then ligated into the vector pAMG21 and transformed into competent E. coli strain 2596 cells as described for Fc-EMP. Clones were screened for ability to produce the recombinant protein product and possession of the gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #3813.

The nucleotide and amino acid sequences (SEQ ID NOS: 19 and 20, respectively) of the resulting fusion protein are shown in Figure 15. There is a silent mutation at position 145 (A to G, shown in boldface) such that the final construct has a different nucleotide sequence than the oligonucleotide 1871-72 from which it was derived.

<u>Fc-EMP-EMP</u>. A DNA sequence coding for the Fc region of human IgG1 fused in-frame to a dimer of the EPO-mimetic peptide was

constructed using standard PCR technology. Templates for PCR reactions were the plasmids from strains 3688 and 3813 above.

The Fc portion of the molecule was generated in a PCR reaction with strain 3688 DNA using the primers 1216-52 and 1798-17 (shown above). The EMP dimer portion of the molecule was the product of a second PCR reaction with strain 3813 DNA using the primers 1798-18 (also shown above) and SEQ ID NO: 418, shown below:

1798-20 CTA ATT GGA TCC TCG AGA TTA ACC CCC TTG TGG CTT ACAT

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The oligonucleotides 1798-17 and 1798-18 contain an overlap of 61 nucleotides, allowing the two genes to be fused together in the correct reading frame by combining the above PCR products in a third reaction using the outside primers, 1216-52 and 1798-20.

The final PCR gene product (the full length fusion gene) was digested with restriction endonucleases XbaI and BamHI, and then ligated into the vector pAMG21 and transformed into competent E. coli strain 2596 cells as described for Fc-EMP. Clones were screened for the ability to produce the recombinant protein product and to possess the gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #3822.

The nucleotide and amino acid sequences (SEQ ID NOS: __ and __, respectively) of the fusion protein are shown in Figure 16.

<u>Characterization of Fc-EMP activity</u>. Characterization was carried out <u>in vivo</u> as follows.

Mice: Normal female BDF1 approximately 10-12 weeks of age.

Bleed schedule: Ten mice per group treated on day 0, two groups started 4 days apart for a total of 20 mice per group. Five mice bled at each time point, mice were bled a maximum of three times a week. Mice were anesthetized with isoflurane and a total volume of 140-160 ml of blood was obtained by puncture of the orbital sinus. Blood was counted

on a Technicon H1E blood analyzer running software for murine blood. Parameters measured were WBC, RBC, HCT, HGB, PLT, NEUT, LYMPH.

Treatments: Mice were either injected subcutaneously for a bolus treatment or implanted with 7 day micro-osmotic pumps for continuous delivery. Subcutaneous injections were delivered in a volume of 0.2 ml. Osmotic pumps were inserted into a subcutaneous incision made in the skin between the scapulae of anesthetized mice. Compounds were diluted in PBS with 0.1% BSA. All experiments included one control group, labeled "carrier" that were treated with this diluent only. The concentration of the test articles in the pumps was adjusted so that the calibrated flow rate from the pumps gave the treatment levels indicated in the graphs.

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Experiments: Various Fc-conjugated EPO mimetic peptides (EMPs) were delivered to mice as a single bolus injection at a dose of $100 \,\mu\text{g/kg}$. Fc-EMPs were delivered to mice in 7-day micro-osmotic pumps. The pumps were not replaced at the end of 7 days. Mice were bled until day 51 when HGB and HCT returned to baseline levels.

Example 4

TNF-a inhibitors

Fc-TNF-α inhibitors. A DNA sequence coding for the Fc region of human IgG1 fused in-frame to a monomer of the TNF-α inhibitory peptide was constructed using standard PCR technology. The Fc and 5 glycine linker portion of the molecule was generated in a PCR reaction with DNA from the Fc-EMP fusion strain #3718 (see Example 3) using the sense primer 1216-52 and the antisense primer 2295-89 (SEQ ID NOS: 1112 and 1113, respectively). The nucleotides encoding the TNF-α inhibitory peptide were provided by the PCR primer 2295-89 shown below:

30 2295-89 CCG CGG ATC CAT TAC GGA CGG TGA CCC AGA GAG GTG TTT TTG TAG

TGC GGC AGG AAG TCA CCA CCA CCT CCA CCT TTA CCC

The oligonucleotide 2295-89 overlaps the glycine linker and Fc portion of the template by 22 nucleotides, with the PCR resulting in the two genes being fused together in the correct reading frame.

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The PCR gene product (the full length fusion gene) was digested with restriction endonucleases NdeI and BamHI, and then ligated into the vector pAMG21 and transformed into competent E.coli strain 2596 cells as described for EMP-Fc herein. Clones were screened for the ability to produce the recombinant protein product and to possess the gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #4544.

The nucleotide and amino acid sequences (SEQ ID NOS: 1055 and 1056) of the fusion protein are shown in Figures 19A and 19B.

<u>TNF- α inhibitor-Fc.</u> A DNA sequence coding for a TNF- α inhibitory peptide fused in-frame to the Fc region of human IgG1 was constructed using standard PCR technology. The template for the PCR reaction was a plasmid containing an unrelated peptide fused via a five glycine linker to Fc. The nucleotides encoding the TNF- α inhibitory peptide were provided by the sense PCR primer 2295-88, with primer 1200-54 serving as the antisense primer (SEQ ID NOS: 1117 and 407, respectively). The primer sequences are shown below:

2295-88 GAA TAA CAT ATG GAC TTC CTG CCG CAC TAC AAA AAC ACC TCT CTG GGT CAC CGT CCG GGT GGA GGC GGT GGG GAC AAA ACT

1200-54 GTT ATT GCT CAG CGG TGG CA

The oligonucleotide 2295-88 overlaps the glycine linker and Fc portion of the template by 24 nucleotides, with the PCR resulting in the two genes being fused together in the correct reading frame.

The PCR gene product (the full length fusion gene) was digested with restriction endonucleases <u>Nde</u>I and <u>Bam</u>HI, and then ligated into the vector pAMG21 and transformed into competent <u>E. coli</u> strain 2596 cells as described for EMP-Fc herein. Clones were screened for the ability to produce the recombinant protein product and to possess the gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #4543.

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The nucleotide and amino acid sequences (SEQ ID NOS: 1057 and 1058) of the fusion protein are shown in Figures 20A and 20B.

Expression in E. coli. Cultures of each of the pAMG21-Fc-fusion constructs in E. coli GM221 were grown at 37 °C in Luria Broth medium containing 50 mg/ml kanamycin. Induction of gene product expression from the luxPR promoter was achieved following the addition of the synthetic autoinducer N-(3-oxohexanoyl)-DL-homoserine lactone to the culture media to a final concentration of 20 ng/ml. Cultures were incubated at 37 °C for a further 3 hours. After 3 hours, the bacterial cultures were examined by microscopy for the presence of inclusion bodies and were then collected by centrifugation. Refractile inclusion bodies were observed in induced cultures indicating that the Fc-fusions were most likely produced in the insoluble fraction in E. coli. Cell pellets were lysed directly by resuspension in Laemmli sample buffer containing 10% β -mercaptoethanol and were analyzed by SDS-PAGE. In each case, an intense coomassie-stained band of the appropriate molecular weight was observed on an SDS-PAGE gel.

Purification of Fc-peptide fusion proteins. Cells are broken in water (1/10) by high pressure homogenization (2 passes at 14,000 PSI) and inclusion bodies are harvested by centrifugation (4200 RPM in J-6B for 1 hour). Inclusion bodies are solubilized in 6M guanidine, 50mM Tris, 8mM DTT, pH 8.7 for 1 hour at a 1/10 ratio. The solubilized mixture is diluted

20 times into 2M urea, 50 mM tris, 160mM arginine, 3mM cysteine, pH 8.5. The mixture is stirred overnight in the cold and then concentrated about 10 fold by ultafiltration. It is then diluted 3 fold with 10mM Tris, 1.5M urea, pH 9. The pH of this mixture is then adjusted to pH 5 with acetic acid. The precipitate is removed by centrifugation and the supernatant is loaded onto a SP-Sepharose Fast Flow column equilibrated in 20mM NaAc, 100 mM NaCl, pH 5 (10mg/ml protein load, room temperature). The protein is eluted from the column using a 20 column volume gradient in the same buffer ranging from 100mM NaCl to 500mM NaCl. The pool from the column is diluted 3 fold and loaded onto a SP-Sepharose HP column in 20mM NaAc, 150mM NaCl, pH 5(10mg/ml protein load, room temperature). The protein is eluted using a 20 column volume gradient in the same buffer ranging from 150mM NaCl to 400mM NaCl. The peak is pooled and filtered.

<u>Characterization of activity of Fc-TNF- α inhibitor and TNF- α inhibitor -Fc. Binding of these peptide fusion proteins to TNF- α can be characterized by BIAcore by methods available to one of ordinary skill in the art who is armed with the teachings of the present specification.</u>

Example 5

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IL-1 Antagonists

Fc-IL-1 antagonist. A DNA sequence coding for the Fc region of human IgG1 fused in-frame to a monomer of an IL-1 antagonist peptide was constructed using standard PCR technology. The Fc and 5 glycine linker portion of the molecule was generated in a PCR reaction with DNA from the Fc-EMP fusion strain #3718 (see Example 3) using the sense primer 1216-52 and the antisense primer 2269-70 (SEQ ID NOS: 1112 and 1118, respectively). The nucleotides encoding the IL-1 antagonist peptide were provided by the PCR primer 2269-70 shown below:

1216-52	AAC ATA AGT ACC TGT AGG ATC G
2269-70	CCG CGG ATC CAT TAC AGC GGC AGA GCG TAC GGC TGC CAG TAA CCC GGG GTC CAT TCG AAA CCA CCA CCT CCA CCT TTA CCC

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The oligonucleotide 2269-70 overlaps the glycine linker and Fc portion of the template by 22 nucleotides, with the PCR resulting in the two genes being fused together in the correct reading frame.

The PCR gene product (the full length fusion gene) was digested with restriction endonucleases <u>Nde</u>I and <u>Bam</u>HI, and then ligated into the vector pAMG21 and transformed into competent <u>E. coli</u> strain 2596 cells as described for EMP-Fc herein. Clones were screened for the ability to produce the recombinant protein product and to possess the gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #4506.

The nucleotide and amino acid sequences (SEQ ID NOS: 1059 and 1060) of the fusion protein are shown in Figures 21A and 21B.

<u>IL-1 antagonist-Fc.</u> A DNA sequence coding for an IL-1 antagonist peptide fused in-frame to the Fc region of human IgG1 was constructed using standard PCR technology. The template for the PCR reaction was a plasmid containing an unrelated peptide fused via a five glycine linker to Fc. The nucleotides encoding the IL-1 antagonist peptide were provided by the sense PCR primer 2269-69, with primer 1200-54 serving as the antisense primer (SEQ ID NOS: 1119 and 407, respectively). The primer sequences are shown below:

30	2269-69	GAA CTG									CAG	CCG	TAC	GCT
	1200-54	GTT	ATT	GCT	CAG	CGG	TGG	CA						

The oligonucleotide 2269-69 overlaps the glycine linker and Fc portion of the template by 24 nucleotides, with the PCR resulting in the two genes being fused together in the correct reading frame.

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The PCR gene product (the full length fusion gene) was digested with restriction endonucleases NdeI and BamHI, and then ligated into the vector pAMG21 and transformed into competent E. coli strain 2596 cells as described for EMP-Fc herein. Clones were screened for the ability to produce the recombinant protein product and to possess the gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #4505.

The nucleotide and amino acid sequences (SEQ ID NOS: 1061 and 1062) of the fusion protein are shown in Figures 22A and 22B. Expression and purification were carried out as in previous examples.

Characterization of Fc-IL-1 antagonist peptide and IL-1 antagonist peptide-Fc activity. IL-1 Receptor Binding competition between IL-1β, IL-1RA and Fc-conjugated IL-1 peptide sequences was carried out using the IGEN system. Reactions contained 0.4 nM biotin-IL-1R + 15 nM IL-1-TAG + 3 uM competitor + 20 ug/ml streptavidin-conjugate beads, where competitors were IL-1RA, Fc-IL-1 antagonist, IL-1 antagonist-Fc). Competition was assayed over a range of competitor concentrations from 3 uM to 1.5 pM. The results are shown in Table C below:

Table C—Results from IL-1 Receptor Binding Competition Assay

		IL-1pep-Fc	Fc-IL-1pep	IL-1ra
5	KI EC50	281.5 530.0	59.58 112.2	1.405 2.645
	95% Confidence	e Intervals		
10	EC50	280.2 to 1002	54.75 to 229.8	1.149 to 6.086
15	KI	148.9 to 532.5	29.08 to 122.1	0.6106 to 3.233
Τ.)	Goodness of F	it		
	IR ²	0.9790	0.9687	0.9602

Example 6

VEGF-Antagonists

Fc-VEGF Antagonist. A DNA sequence coding for the Fc region of human IgG1 fused in-frame to a monomer of the VEGF mimetic peptide was constructed using standard PCR technology. The templates for the PCR reaction were the pFc-A3 plasmid and a synthetic VEGF mimetic peptide gene. The synthetic gene was assembled by annealing the following two oligonucleotides primer (SEQ ID NOS: 1120 and 1121, respectively):

2293-11 GTT GAA CCG AAC TGT GAC ATC CAT GTT ATG TGG GAA TGG GAA TGT TTT GAA CGT CTG

2293-12 CAG ACG TTC AAA ACA TTC CCA TTC CCA CAT AAC ATG GAT GTC ACA GTT CGG TTC AAC

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The two oligonucleotides anneal to form the following duplex encoding an amino acid sequence shown below (SEQ ID NOS 1122):

This duplex was amplified in a PCR reaction using 2293-05 and 2293-06 as the sense and antisense primers (SEQ ID NOS. 1125 and 1126).

The Fc portion of the molecule was generated in a PCR reaction with the pFc-A3 plasmid using the primers 2293-03 and 2293-04 as the sense and antisense primers (SEQ ID NOS. 1123 and 1124, respectively). The full length fusion gene was obtained from a third PCR reaction using the outside primers 2293-03 and 2293-06. These primers are shown below:

	2293-03			TTC	TAG	AAG	GAG	GAA	TAA	CAT	ATG	GAC	AAA	ACT	CAC
		ACA	TGT												
5	2293-04	GTC	ACA	GTT	CGG	TTC	AAC	ACC	ACC	ACC	ACC	ACC	TTT	ACC	CGG
		AGA	CAG	GGA											
	2293-05	TCC	CTG	TCT	CCG	GGT	AAA	GGT	GGT	GGT	GGT	GGT	GTT	GAA	CCG
		AAC	TGT	GAC	ATC										
10															
	2293-06	CCG	CGG	ATC	CTC	GAG	TTA	CAG	ACG	TTC	AAA	ACA	TTC	CCA	

The PCR gene product (the full length fusion gene) was digested with restriction endonucleases <u>NdeI</u> and <u>BamHI</u>, and then ligated into the vector pAMG21 and transformed into competent <u>E. coli</u> strain 2596 cells as described for EMP-Fc herein. Clones were screened for the ability to produce the recombinant protein product and to possess the gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #4523.

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The nucleotide and amino acid sequences (SEQ ID NOS: 1063 and 1064) of the fusion protein are shown in Figures 23A and 23B.

<u>VEGF antagonist -Fc</u>. A DNA sequence coding for a VEGF mimetic peptide fused in-frame to the Fc region of human IgG1 was constructed using standard PCR technology. The templates for the PCR reaction were the pFc-A3 plasmid and the synthetic VEGF mimetic peptide gene described above. The synthetic duplex was amplified in a PCR reaction using 2293-07 and 2293-08 as the sense and antisense primers (SEQ ID NOS. 1127 and 1128, respectively).

The Fc portion of the molecule was generated in a PCR reaction with the pFc-A3 plasmid using the primers 2293-09 and 2293-10 as the sense and antisense primers (SEQ ID NOS. 1129 and 1130, respectively).

The full length fusion gene was obtained from a third PCR reaction using the outside primers 2293-07 and 2293-10. These primers are shown below:

	2293-07	ATT	TGA	TTC	TAG	AAG	GAG	GAA	TAA	CAT	ATG	GTT	GAA	CCG	AAC
5		TGT	GAC												
	2293-08	ACA	TGT	GTG	AGT	TTT	GTC	ACC	ACC	ACC	ACC	ACC	CAG	ACG	TTC
		AAA	ACA	TTC											
10	2293-09		TGT ACA		GAA	CGT	CTG	GGT	GGT	GGT	GGT	GGT	GAC	AAA	ACT

The PCR gene product (the full length fusion gene) was digested
with restriction endonucleases NdeI and BamHI, and then ligated into the vector pAMG21 and transformed into competent E. coli strain 2596 cells as described for EMP-Fc herein. Clones were screened for the ability to produce the recombinant protein product and to possess the gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #4524.

The nucleotide and amino acid sequences (SEQ ID NOS: 1065 and 1066) of the fusion protein are shown in Figures 24A and 24B. Expression and purification were carried out as in previous examples.

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Example 7 MMP Inhibitors

Fc-MMP inhibitor. A DNA sequence coding for the Fc region of human IgG1 fused in-frame to a monomer of an MMP inhibitory peptide was constructed using standard PCR technology. The Fc and 5 glycine linker portion of the molecule was generated in a PCR reaction with DNA from the Fc-TNF- α inhibitor fusion strain #4544 (see Example 4) using the sense primer 1216-52 and the antisense primer 2308-67 (SEQ ID NOS: 1112

and 1131, respectively). The nucleotides encoding the MMP inhibitor peptide were provided by the PCR primer 2308-67 shown below:

The oligonucleotide 2308-67 overlaps the glycine linker and Fc portion of the template by 22 nucleotides, with the PCR resulting in the two genes being fused together in the correct reading frame.

The PCR gene product (the full length fusion gene) was digested with restriction endonucleases <u>Ndel</u> and <u>BamHI</u>, and then ligated into the vector pAMG21 and transformed into competent <u>E. coli</u> strain 2596 cells as described for EMP-Fc herein. Clones were screened for the ability to produce the recombinant protein product and to possess the gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #4597.

The nucleotide and amino acid sequences (SEQ ID NOS: 1067 and 1068) of the fusion protein are shown in Figures 25A and 25B. Expression and purification were carried out as in previous examples.

MMP Inhibitor-Fc. A DNA sequence coding for an MMP inhibitory peptide fused in-frame to the Fc region of human IgG1 was constructed using standard PCR technology. The Fc and 5 glycine linker portion of the molecule was generated in a PCR reaction with DNA from the Fc-TNF- α inhibitor fusion strain #4543 (see Example 4). The nucleotides encoding the MMP inhibitory peptide were provided by the sense PCR primer 2308-66, with primer 1200-54 serving as the antisense primer (SEQ ID NOS: 1132 and 407, respectively). The primer sequences are shown below:

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2308-66 GAA TAA CAT ATG TGC ACC CAC TGG GGT TTC ACC CTG TGC GGT GGA GGC GGT GGG GAC AAA

35 1200-54 GTT ATT GCT CAG CGG TGG CA

The oligonucleotide 2269-69 overlaps the glycine linker and Fc portion of the template by 24 nucleotides, with the PCR resulting in the two genes being fused together in the correct reading frame.

The PCR gene product (the full length fusion gene) was digested with restriction endonucleases Ndel and BamHI, and then ligated into the vector pAMG21 and transformed into competent E. coli strain 2596 cells as described for EMP-Fc herein. Clones were screened for the ability to produce the recombinant protein product and to possess the gene fusion having the correct nucleotide sequence. A single such clone was selected and designated Amgen strain #4598.

The nucleotide and amino acid sequences (SEQ ID NOS: 1069 and 1070) of the fusion protein are shown in Figures 26A and 26B.

The invention now being fully described, it will be apparent to one of ordinary skill in the art that many changes and modifications can be made thereto, without departing from the spirit and scope of the invention as set forth herein.

20 Abbreviations

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Abbreviations used throughout this specification are as defined below, unless otherwise defined in specific circumstances.

٠	Ac T	acetyl (used to refer to acetylated residues)
	AcBpa	acetylated p-benzoyl-L-phenylalanine
25	ADCC	antibody-dependent cellular cytotoxicity
	Aib	aminoisobutyric acid
	··· bA	beta-alarine
	Вра	p-benzoyl-L-phenylalanine
	BrAc	bromoacetyl (BrCH2C(O)

	BSA	Bovine serum albumin
	Bzl	Benzyl
	Cap	Caproic acid
	CTL	Cytotoxic T lymphocytes
5	CTLA4	Cytotoxic T lymphocyte antigen 4
	DARC	Duffy blood group antigen receptor
	DCC	Dicylcohexylcarbodiimide
	Dde	1-(4,4-dimethyl-2,6-dioxo-cyclohexylidene)ethyl
	EMP	Erythropoietin-mimetic peptide
10	ESI-MS	Electron spray ionization mass spectrometry
	EPO	Erythropoietin
	Fmoc	fluorenylmethoxycarbonyl
	G-CSF	Granulocyte colony stimulating factor
	GH	Growth hormone
15	HCT	hematocrit
•	HGB	hemoglobin
	hGH	Human growth hormone
	HOBt	1-Hydroxybenzotriazole
	HPLC	high performance liquid chromatography
20	IL	interleukin
	IL-R	interleukin receptor
	IL-1R	interleukin-1 receptor
	IL-1ra	interleukin-1 receptor antagonist
	Lau	Lauric acid
25	LPS	lipopolysaccharide
	LYMPH	lymphocytes
•••	MALDI-MS	Matrix-assisted laser desorption ionization mass
		spectrometry
	Me	methyl

	MeO	methoxy
	MHC	major histocompatibility complex
	MMP	matrix metalloproteinase
	MMPI	matrix metalloproteinase inhibitor
5	1-Nap	1-napthylalanine
	NEUT	neutrophils
	NGF	nerve growth factor
	Nle	norleucine
	NMP	N-methyl-2-pyrrolidinone
10	PAGE	polyacrylamide gel electrophoresis
	PBS	Phosphate-buffered saline
	Pbf	2,2,4,6,7-pendamethyldihydrobenzofuran-5-sulfonyl
	PCR	polymerase chain reaction
	Pec	pipecolic acid
15	PEG	Poly(ethylene glycol)
	pGlu	pyroglutamic acid
	Pic	picolinic acid
	PLT	platelets
	pΥ	phosphotyrosine
20	RBC	red blood cells
	RBS	ribosome binding site
	RT	room temperature (25 °C)
	Sar	sarcosine
	SDS	sodium dodecyl sulfate
25	STK	serine-threonine kinases
	t-Boc	tert-Butoxycarbonyl
***	tBu	tert-Butyl
	TGF	tissue growth factor
	THF	thymic humoral factor

TK tyrosine kinase TMP Thrombopoietin-mimetic peptide TNF Tissue necrosis factor TPO Thrombopoietin TNF-related apoptosis-inducing ligand 5 TRAIL Trt trityl UK urokinase UKR urokinase receptor vascular endothelial cell growth factor **VEGF** VIP vasoactive intestinal peptide 10 **WBC** white blood cells

What is claimed is:

1. A composition of matter of the formula

$$(X^1)_a - F^1 - (X^2)_b$$

and multimers thereof, wherein:

5 F¹ is an Fc domain;

 X^{1} and X^{2} are each independently selected from - $(L^{1})_{c}$ - P^{1} , - $(L^{1})_{c}$ - P^{1} - $(L^{2})_{d}$ - P^{2} - $(L^{1})_{c}$ - P^{1} - $(L^{2})_{d}$ - P^{2} - $(L^{3})_{e}$ - P^{3} , and - $(L^{1})_{c}$ - P^{1} - $(L^{2})_{d}$ - P^{2} - $(L^{3})_{e}$ - P^{3} - $(L^{4})_{c}$ - P^{4}

P¹, P², P³, and P⁴ are each independently sequences of pharmacologically active peptides;

L¹, L², L³, and L⁴ are each independently linkers; and a, b, c, d, e, and f are each independently 0 or 1, provided that at least one of a and b is 1.

2. The composition of matter of Claim 1 of the formulae

15 X'-F'

or

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 F^1-X^2 .

- 3. The composition of matter of Claim 1 of the formula F^1 - (L^1) - P^1 .
- 20 4. The composition of matter of Claim 1 of the formula $F^1\text{-}(L^1)_c\text{-}P^1\text{-}(L^2)_d\text{-}P^2.$
 - 5. The composition of matter of Claim 1 wherein F¹ is an IgG Fc domain.
 - 6. The composition of matter of Claim 1 wherein F¹ is an IgG1 Fc domain.
 - 7. The composition of matter of Claim 1 wherein F¹ comprises the sequence of SEQ ID NO: 2.
 - 8. The composition of matter of Claim 1 wherein X¹ and X² comprise an IL-1 antagonist peptide sequence.

9. The composition of matter of Claim 8 wherein the IL-1 antagonist peptide sequence is selected from SEQ ID NOS: 212, 907, 908, 909, 910, 917, and 979.

10. The composition of matter of Claim 8 wherein the IL-1 antagonist peptide sequence is selected from SEQ ID NOS: 213 to 271, 671 to 906, 911 to 916, and 918 to 1023.

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- 11. The composition of matter of Claim 8 wherein F¹ comprises the sequence of SEQ ID NO: 2.
- The composition of matter of Claim 1 wherein X¹ and X² comprise
 an EPO-mimetic peptide sequence.
 - 13. The composition of matter of Claim 12 wherein the EPO-mimetic peptide sequence is selected from Table 5.
 - 14. The composition of matter of Claim 12 wherein F¹ comprises the sequence of SEQ ID NO: 2.
- 15 15. The composition of matter of Claim 12 comprising a sequence selected from SEQ ID NOS: 83, 84, 85, 124, 419, 420, 421, and 461.
 - 16. The composition of matter of claim 12 comprising a sequence selected from SEQ ID NOS: 339 and 340.
- The composition of matter of Claim 12 comprising a sequence
 selected from SEQ ID NOS: 20 and 22.
 - 18. The composition of matter of Claim 3 wherein P¹ is a TPO-mimetic peptide sequence.
 - 19. The composition of matter of Claim 18 wherein P¹ is a TPO-mimetic peptide sequence selected from Table 6.
- 25 20. The composition of matter of Claim 18 wherein F¹ comprises the sequence of SEQ ID NO: 2.
 - 21. The composition of matter of Claim 18 having a sequence selected from SEQ ID NOS: 6 and 12.
 - 22. A DNA encoding a composition of matter of any of Claims 1 to 21.

- 23. An expression vector comprising the DNA of Claim 22.
- 24. A host cell comprising the expression vector of Claim 23.
- 25. The cell of Claim 24, wherein the cell is an <u>E. coli</u> cell.

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- 26. A process for preparing a pharmacologically active compound, which comprises
 - selecting at least one randomized peptide that modulates the
 activity of a protein of interest; and
 - b) preparing a pharmacologic agent comprising at least one Fc domain covalently linked to at least one amino acid sequence of the selected peptide or peptides.
- 27. The process of Claim 26, wherein the peptide is selected in a process comprising screening of a phage display library, an <u>E. coli</u> display library, a ribosomal library, or a chemical peptide library.
- 28. The process of Claim 26, wherein the preparation of the pharmacologic agent is carried out by:
 - a) preparing a gene construct comprising a nucleic acid
 sequence encoding the selected peptide and a nucleic acid
 sequence encoding an Fc domain; and
 - b) expressing the gene construct.
- 20 29. The process of Claim 26, wherein the gene construct is expressed in an <u>E. coli</u> cell.
 - 30. The process of Claim 26, wherein the protein of interest is a cell surface receptor.
- 31. The process of Claim 26, wherein the protein of interest has a linear epitope.
 - 32. The process of Claim 26, wherein the protein of interest is a cytokine receptor.
 - 33. The process of Claim 26, wherein the peptide is an EPO-mimetic peptide.

34. The process of Claim 26, wherein the peptide is a TPO-mimetic peptide.

- 35. The process of Claim 26, wherein the peptide is an IL-1 antagonist peptide.
- 5 36. The process of Claim 26, wherein the peptide is an MMP inhibitor peptide or a VEGF antagonist peptide.
 - 37. The process of Claim 26, wherein the peptide is a TNF-antagonist peptide.
- 38. The process of Claim 26, wherein the peptide is a CTLA4-mimetic peptide.
 - 39. The process of Claim 26, wherein the peptide is selected from Tables 4 to 20.
 - 40. The process of Claim 26, wherein the selection of the peptide is carried out by a process comprising:
- a) preparing a gene construct comprising a nucleic acid sequence encoding a first selected peptide and a nucleic acid sequence encoding an Fc domain;

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- b) conducting a polymerase chain reaction using the gene construct and mutagenic primers, wherein
 - i) a first mutagenic primer comprises a nucleic acid sequence complementary to a sequence at or near the
 5' end of a coding strand of the gene construct, and
 - ii) a second mutagenic primer comprises a nucleic acid sequence complementary to the 3' end of the noncoding strand of the gene construct.
- 41. The process of Claim 26, wherein the compound is derivatized.
- 42. The process of Claim 26, wherein the derivatized compound comprises a cyclic portion, a cross-linking site, a non-peptidyl

linkage, an N-terminal replacement, a C-terminal replacement, or a modified amino acid moiety.

- 43. The process of Claim 26 wherein the Fc domain is an IgG Fc domain.
- 5 44. The process of Claim 26, wherein the vehicle is an IgG1 Fc domain.
 - 45. The process of Claim 26, wherein the vehicle comprises the sequence of SEQ ID NO: 2.
 - 46. The process of Claim 26, wherein the compound prepared is of the formula

 $(X^1)_a - F^1 - (X^2)_b$

and multimers thereof, wherein:

F¹ is an Fc domain;

 X^{1} and X^{2} are each independently selected from $-(L^{1})_{c} - P^{1}$, $-(L^{1})_{c} - P^{1} - (L^{2})_{d} - P^{2}$, $-(L^{1})_{c} - P^{1} - (L^{2})_{d} - P^{2} - (L^{3})_{e} - P^{3}$, and $-(L^{1})_{c} - P^{1} - (L^{2})_{d} - P^{2} - (L^{3})_{e} - P^{3} - (L^{4})_{c} - P^{4}$

P¹, P², P³, and P⁴ are each independently sequences of pharmacologically active peptides;

L¹, L², L³, and L⁴ are each independently linkers; and a, b, c, d, e, and f are each independently 0 or 1, provided that at least one of a and b is 1.

47. The process of Claim 46, wherein the compound prepared is of the formulae

Y1_F1

or

15

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F¹-X².

48. The process of Claim 46, wherein the compound prepared is of the formulae

 \mathbf{or}

$$F^{1}-(L^{1})_{c}-P^{1}-(L^{2})_{d}-P^{2}.$$

- 49. The process of Claim 46, wherein F¹ is an IgG Fc domain.
- 50. The process of Claim 46, wherein F¹ is an IgG1 Fc domain.
- 5 51. The process of Claim 46, wherein F¹ comprises the sequence of SEQ ID NO: 2.

peptide selection

peptide optimization

formation of Fc-peptide DNA construct

insertion of construct into expression vector

transfection of host cell with vector

expression of vector in host cell

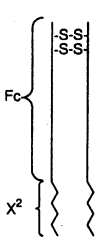
Fc multimer formation in host cell

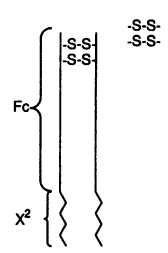
isolation of Fc multimer from host cell

FIG. 2A

FIG. 2B

FIG. 2C





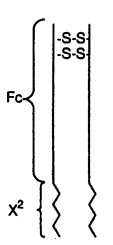
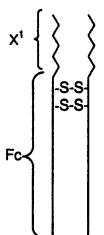
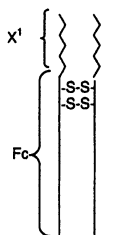


FIG. 2D FIG. 2E

FIG. 2F





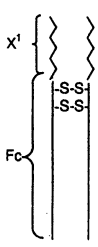


FIG. 3A

FIG. 3B

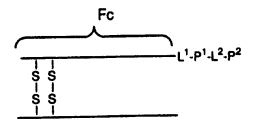
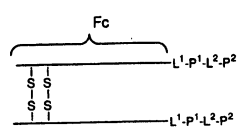


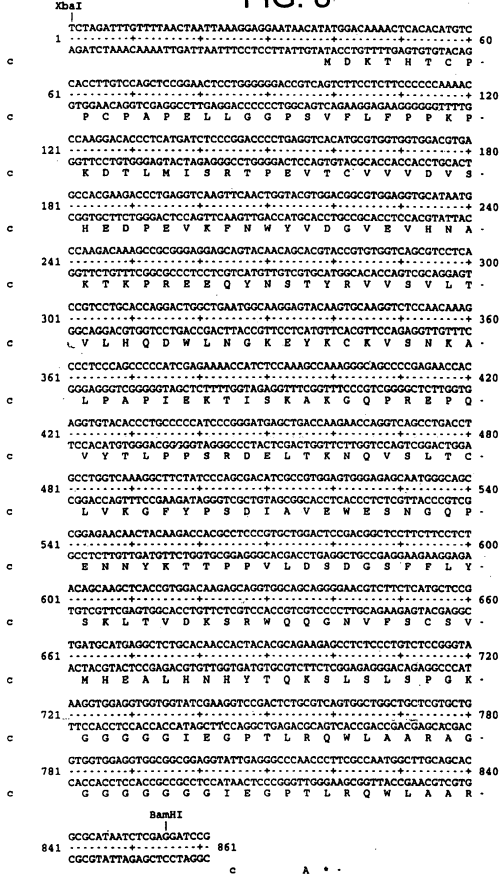
FIG. 3C



	1	ATGGACAAAACTCACATGTCCACCTTGTCCAGCTCCGGAACTCCTGGGGGGACCGTCA													60						
	^	TACC	TGTT	TTG	AGT	GTG?	CACA								GAG	GAC	ccc	CCI	GGC	AGT	00
a		M C	K	T	H	T	С	P	P	С	P	A	P	E	L	L	G	G	P	S	-
	61		TCCT	CTT	CCC	CCC					ACC			ATC					GAG		120
	0.1		LAGGA	GAA(GGG	GG 2						•							CTC		120
a		V F	L	F	P	P	K	P	K	D	T	L	M	I	S	R	T	P	E	V	•
	121		GCGT																		180
	121		CGCA																		100
a		T C	v	v	v	D	V	s	H	E	D	P	E	V	K	F	N	W	Y	V	•
•	101	GAC	GCGT	GGA	GGT	GCA?	raan										TAC	AAC	AGC	ACG	240
	181	CTGC	CGCA	CCT	CCA	CGT	\TT!							CTC			ATG	TTC	TC	TGC	240
a		ם מ	v	E	v	н	N	A	ĸ	T	K	P	R	E	E	Q	Y	N	S	T	-
		TAC	GTGT																GAG	TAC	200
	241	ATG	CACA	CCA			•				•			•			-		CTC	ATG	300
a		Y. I	v s	v	s	v	L	T	v	L	H	Q	D	W	L	N	G	ĸ	E	Y	•
			rgcaa																		260
	301		ACGTI																		360
a		K (C K	v	s	N	ĸ	A	L	P	A	P	I	E	K	Ť	Ï	S	ĸ	A	-
			GGCA	'GCC	CCG	AGA	ACC	ACA	GT(GAT	GAG	CTC	ACC	420
	361		CCGI	CGG	GGC'	TCT	rgg	rgt	CCA					GG1			CTA	CTC	GAC	TGG	420
a		K (G Q	P	R	E	P	Q	v	Y	T	L	P	P	s	R	D	E	L	T	•
	401		AACCA		CAG	CCT															480
	421		rtggi		GTC	GGA(+ GTT				AGGG						400
a		K I	9 Q	v	s	L	T	С	L	v	K	G	F	Y	P	S	D	I	A	V	-
	401	GAG:	rggg <i>i</i>	GAG	CAA	TGG	GCA	GCC	GGA(GAA	CAA	CTA	CAA	GAC	CAC	CC1	CCC	GTO	CTC	GAC	540
	401	CTC	ACCCT	CTC	GTT.	ACC	CGT	CGG	CCT	CTT	GTT(GAT(GTT(CTG	GTGC	:GG/	\GGC	CAC	GAC	CTG	240
a		E 1	W E	s	N	G	Q	P	E	N	N	Y	K	T	T	P	P	٧	L	D	-
		TCC	GACGO	CTC	CTT	CTT	CCT	CTA	CAG	CAA	GCT(CAC	CGT	GGA(CAAC	GAG	AGC	TG	CAC	CAG	600
	241	AGG	CTGCC	GAG	GAA	GAA	GGA	GAT(GTC	GTT	CGA	GTG	GCA	CTC	GTT(CTC	HC(CAC	GTC	GTC	200
a		s i	D G	s	F	F	L	Y	S	K	L	T	V	D	K	S	R	W	Q	Q	-
	CO1	GGG	AACG'	CTT	CTC	ATG	CTC	CGT	GAT	GCA	TGA	GGC'	TCT	GCA	CAA	CA	TAC	CAC	CA	BAAG	660
	901	CCC'	TTGC	GAA	GAG	TAC	GAG	GCA	CTA	CGT	ACT	CCG	AGA	CGT	GTT(GGT(ATC	STG	CGT	TTC	550
a		G I	N V	F	S	С	S	V	M	H	E	A	L	Н	N	H	Y	T	Q	K	•
	661		CTCT							684											
	001		GAGA							J J 4											

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		XbaI											J.	1							
	1	TCTAC												TAT							60
С	-	AGATO														GTT	TTG		GTG'	TACAC	
	61			+				 -			-+-	• • •		+				+			120
c		GTGG/ P												TCA V							;
	121	CCAA		+	·	• • • ·	4	 -			•+•	• • •		+	• • •			+		••••	- 180
С		GGTT(GCACT V S	
	181	GCCA																			
c		CGGT(
	241	CCAA																			300
c		GGTT(CAT Y	-			GTC(S			
	301	CCGT																			360
c		GGCAC V	GA(GTTTC K /	
	361	CCCT																			
C	302	GGGAG	GG1	rcgo	GGG	STAC	CTC	TT	MG	STA	GAG	GTT	TCG	GTT	TCC	CGT	CGG	GGC'	rct'		;
	421	AGGT																			
c	701	TCCAC	CATO	TG	GAC	CGGC	GGT	PAGC	GCC	CT	ACT	CGA	CTG		CTT	GGT	CCA	GTC	GGA(CTGG	
	481	GCCT												GGA							540
c	101	CGGAC		TT	rccc	BAAC	LTA:	\GG(STC	GCT(GTA	GCG	GCA		ÇAC				ACC	CGTC	
	541			• • •	• • • •		• +		• • •	• • •	-+-			+	• • •		• • •	+	- • •	+	600
c		GCCTC	OTTC N	otto N	GAT(Y	STTC K	TGC T	TGC T	P P	AGG:	GCA V	CGA L	CCI D	'GAG S	GCT D	GCC G	GAG S			GGAGA L	
	601	ACAGO																			660
c		TGTC	STT(K	CGA(T T	CAC V	D	TTC K	STC	STC R	CAC W	CGT	CG1 Q	CCC G	CTT N	GCA V	gaa F	GAG'	rac(GAGG(S \	
	661	TGAT	CAT	rga(GC1	CTC	CAC	CAAC	CAC	CTA	CAC	GCA	GAA	GAG	CCT	CTC	CCT	GTC:	rcc	GGGT	- 720
С	001	ACTAC	CGT	CTO	CCG	AGAC	CGTC	TTC	GT(GAT(GTG	CGT	CTI	CTC	GGA	GAG	GGA	CAG	AGG	G I	•
	721	AAGG		+ .			4	 -			-+-			+	• • •			+		1	780
c		TTCC	ACC?	CCI	ACC	ACCA	ATA	CT	rcc/	AGG	CTG	AGA	CGC	AGT Q	CAC	CGA	CCG.	ACG	AGC.	ACGA	
		1	Bami	II 																	
	781	AATC1					4														
		TTAG					-														



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XbaI

FIG. 9

		TCTAGATTTGTTTTAACTAATTAAAGGAGGAATAACATATGATCGAAGGTCCGACTCTGC	
c	1	AGATCTAAACAAAATTGATTAATTTCCTCCTTATTGTATACTAGCTTCCAGGCTGAGACG M I E G P T L R	
	61	GTCAGTGGCTGGCTGGTGGTGGCGGTGGCGAGGGGGTGGCATTGAGGGCCCAA CAGTCACCGACGACGACGACGACGACCACCACCGCCTCCCCACCGTAACTCCCGGGTT	120
c		Q W L A A R A G G G G G G G I E G P T - CCCTTCGCCAATGGCTTGCAGCACGCGCAGGGGGGGGGG	-
C	121		180
	181	GTCCACCTTGCCCAGCACCTGAACTCCTGGGGGGACCGTCAGTTTTCCTCTTCCCCCCAA	240
c		CAGGTGGAACGGGTCGTGGACTTGAGGACCCCCCTGGCAGTCAAAAGGAGAAGGGGGGTT PPCPAPELLGGPSVFLFPPK	•
	241	AACCCAAGGACACCCTCATGATCTCCCGGACCCCTGAGGTCACATGCGTGGTGGACG TTGGGTTCCTGTGGGAGTACTAGAGGGCCTGGGGACTCCAGTGTACGCACCACCACCACCTGC P K D T L M I S R T P E V T C V V V D V	
c	201	TGAGCCACGAAGACCCTGAGGTCAAGTTCAACTGGTACGTGGACGGCGTGGAGGTGCATA	
c	201	ACTCGGTGCTTCTGGGACTCCAGTTCAAGTTGACCATGCACCTGCCGCACCTCCACGTAT S H E D P E V K F N W Y V D G V E V H N	
	361	ATGCCAAGACAAAGCCGCGGGAGGAGCAGTACAACAGCACGTACCGTGTCGTCACCGTCCCTCGTCATGTTGTCGTCATGGCACACCAGTCGCAGG	420
С		A K T K P R E E Q Y N S T Y R V V S V L TCACCGTCCTGCACCAGGACTGGCTGAATGGCAAGGAGTACAAGTGCAAGGTCTCCAACA	
c	421	AGTGGCAGGACGTGGTCCTGACCGACTTACCGTTCCTCATGTTCACGTTCCAGAGGTTGT T V L H Q D W L N G K E Y K C K V S N K	
	481	AAGCCCTCCCAGCCCCCATCGAGAAAACCATCTCCAAAGCCAAAGGGCAGCCCCGAGAAC TTCGGGAGGGTCGGGGGTAGCTCTTTTGGTAGAGGTTTCGGTTTCCCGTCGGGGCTCTTTG A L P A P I E K T I S K A K G Q P R E P	
С	541	CACAGGTGTACACCCTGCCCCCATCCCGGGATGAGCTGACCAAGAACCAGGTCAGCCTGA	600
с		GTGTCCACATGTGGGACGGGGGTAGGGCCCTACTCGACTGGTTCTTGGTCCAGTCGGACT Q V Y T L P P S R D E L T K N Q V S L T	•
c	601	CCTGCCTGGTCAAAGGCTTCTATCCCAGCGACATCGCCGTGGAGTGGGAGAGCAATGGGC GGACGGACCAGTTTCCGAAGATAGGGTCGCTGTAGCGGCACCTCACCCTCTCGTTACCCG C L V K G F Y P S D I A V E W E S N G Q	
Ĭ	661	AGCCGGAGAACAACTACAAGACCACGCCTCCCGTGCTGGACTCCGACGGCTCCTTCTTCC	
С		TCGGCCTCTTGTTGATGTTCTGGTGCGGAGGGCACGACCTGAGGCTGCCGAGGAAGAAGG PENNYKTTPPVLDSDGSFFL	•
c	721	TCTACAGCAAGCTCACCGTGGACAAGAGCAGGTGGCAGCAGGGGAACGTCTTCTCATGCT AGATGTCGTTCGAGTGGCACCTGTTCTCGTCCACCGTCGTCCCCTTGCAGAAGAGTACGA Y S K L T V D K S R W Q Q G N V F S C S	
	781	CCGTGATGCATGAGGCTCTGCACAACCACTACACGCAGAAGAGCCTCTCCCTGTCTCCGG GGCACTACGTACTCCGAGACGTGTTGGTGATGTGCGTCTTCTCGGAGAGGGACAGAGGCC V M H Z A L H N H Y T Q K S L S L S P G	
С		BamHI	
		GTAAATAATGGATCC	
	841	CATTTATTACCTAGG	

FIG. 10

	3	baI IIO. IO
	1	 TCTAGATTTGTTTTAACTAATTAAAGGAGGAATAACATATGATCGAAGGTCCGACTCTGC +++
С		M I E G P T L R -
c	61	CAGTCACCGACGAGCACGACCACCTCCGCCACCCCTGTTTTGAGTGTGTACAGGTG Q W L A A R A G G G G D K T H T C P P
c	121	CTTGCCCAGCACCTGAACTCCTGGGGGGGACCGTCAGTTTTCCTCTTCCCCCCAAAACCCA + 180 GAACGGGTCGTGGACTTGAGGACCCCCCTGGCAGTCAAAAGGAGAGGGGGGTTTTGGGT C P A P E L L G G P S V F L F P P K P K
	181	AGGACACCCTCATGATCTCCCGGACCCCTGAGGTCACATGCGTGGTGGTGGACGTGAGCC 240 TCCTGTGGGAGTACTAGAGGGCCTGGGGACTCCAGTGTACGCACCACCACCACCTGCACTCGG
С	241	D T L M I S R T P E V T C V V D V S H - ACGAAGACCCTGAGGTCAAGTTCAACTGGTACGTGGACGCGTGGAGGTGCATAATGCCA
С	201	E D P E V R F N W Y V D G V E V H N A R - AGACAAAGCCGCGGGAGGAGCAGTACAACAGCACGTACCGTGTGGTCAGCGTCCTCACCG
c	301	TCTGTTTCGGCGCCCTCCTCGTCATGTTGTCGTGCATGGCACCAGTCGCAGGAGTGGC T R P R E E Q Y N S T Y R V V S V L T V -
c	361	AGGACGTGGTCCTGACCGACTTACCGTTCCTCATGTTCACGTTCCAGAGGTTGTTTCGGG L H Q D W L N G K E Y K C K V S N K A L -
c	421	TCCCAGCCCCATCGAGAAAACCATCTCCAAAGCCAAAGGGCAGCCCCGAGAACCACAGG
c	481	TGTACACCCTGCCCCATCCCGGGATGAGCTGACCAAGAACCAGGTCAGCCTGACCTGCC ACATGTGGGACGGGGTAGGGCCCTACTCGACTGGTTCTTGGTCCAGTCGGACTGACGG Y T L P P S R D E L T K N Q V S L T C L
Č	541	TGGTCAAAGGCTTCTATCCCAGCGACATCGCCGTGGAGTGGGAGAGCAATGGGCAGCCGG
С	601	
С		TCTTGTTGATGTTCTGGTGCGGAGGGCACGACCTGAGGCTGCCGAGGAAGAAGGAGATGT N N Y K T T P P V L D S D G S F F L Y S GCAAGCTCACCGTGGACAAGAGCAGGTGGCAGCAGGGGAACGTCTTCTCATGCTCCGTGA
c		CGTTCGAGTGGCACCTGTTCTCGTCCACCGTCGTCCCCTTGCAGAAGAGTACGAGGCACT K L T V D K S R W Q Q G N V F S C S V M
c	721	TGCATGAGGCTCTGCACAACCACTACACGCAGAAGAGGCCTCTCCCTGTCTCCGGGTAAAT 780 ACGTACTCCGAGACGTGTTGGTGATGTGCGTCTTCTCGGAGAGGGACAGAGGCCCATTTA H E A L H N H Y T Q K S L S L S P G K *
	781	BamHI AATGGATCC 789 TTACCTAGG

FIG.11

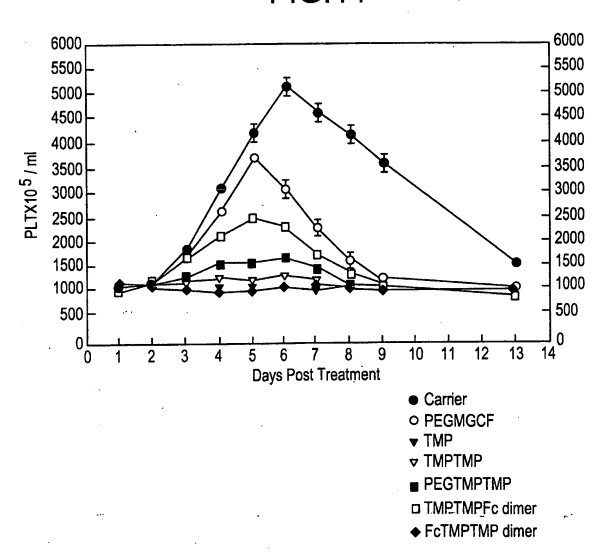
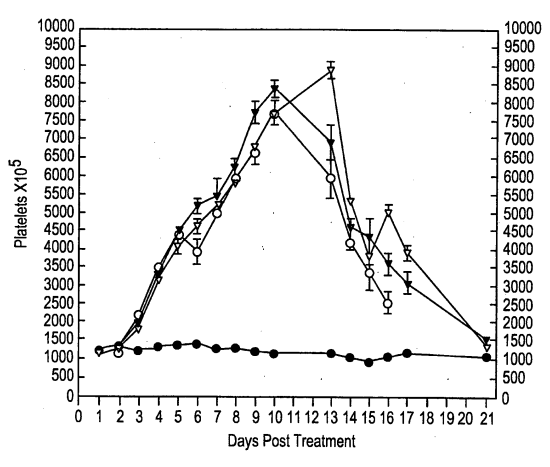


FIG.12



- Carrier
- O PEG MGDF
- ▼ TMPTMPFc dimer
- ▼ FcTMPTMP dimer

FIG. 13

	•	xDa1	
		TCTAGATTTGTTTTAACTAATTAAAGGAGGAATAACATATGGACAAAACTCACACATGTC	
	1		60
c		AGATCTAAACAAAATTGATTAATTTCCTCCTTATTGTATACCTGTTTTGAGTGTGTACAG M D K T H T C P	
•		CACCTTGTCCAGCTCCGGAACTCCTGGGGGGACCGTCAGTCTTCCTCTTCCCCCCAAAAC	
	61		120
c		GTGGAACAGGTCGAGGCCTTGAGGACCCCCCTGGCAGTCAGAAGGAGGAGGAGGGGGTTTTG PCPAPELLGGPSVFLFPPKP	
Ç		r C r R r E D D G t D Y t D t r r R r	
		CCAAGGACACCCTCATGATCTCCCGGACCCCTGAGGTCACATGCGTGGTGGTGGACGTGA	
	121	GGTTCTGTGGGAGTACTAGAGGGCCTGGGGACTCCAGTGTACGCACCACCACCTGCACT	180
С		K D T L M I S R T P E V T C V V V D V S	
•			
		GCCACGAAGACCCTGAGGTCAAGTTCAACTGGTACGTGGACGCCTTGGAGGTGCATAATG	242
	181	CGGTGCTTCTGGGACTCCAGTTCAAGTTGACCATGCACCTGCCGCACCTCCACGTATTAC	240
c		H E D P E V K F N W Y V D G V E V H N A	
_	•		
		CCAAGACAAAGCCGCGGGAGGAGCAGTACAACAGCACGTACCGTGTGGTCAGCGTCCTCA	200
	241	GGTTCTGTTTCGGCGCCCTCCTCGTCATGTTGTCGTGCATGGCACACCAGTCGCAGGAGT	300
С		KTKPREEQYNSTYRVVSVLT	-
	201	CCGTCCTGCACCAGGACTGCCTGAATGGCAAGGAGTACAAGTGCAAGGTCTCCAACAAAG	360
	301	GGCAGGACGTGGTCCTGACCGACTTACCGTTCCTCATGTTCACGTTCCAGAGGTTGTTTC	300
С		V L H Q D W L N G K E Y K C K V S N K A	•
•			
	CC	CTCCCAGCCCCCATCGAGAAAACCATCTCCAAAGCCAAAGGGCAGCCCCGAGAACCAC	
	361		420
_		GGGAGGGTCGGGGTAGCTCTTTTGGTAGAGGTTTCGGTTTCCCGTCGGGGCTCTTGGTG L P A P I E K T I S K A K G Q P R E P Q	
C			
		AGGTGTACACCCTGCCCCCATCCCGGGATGAGCTGACCAAGAACCAGGTCAGCCTGACCT	
	421	TCCACATGTGGGACGGGGTAGGGCCCTACTCGACTGGTTCTTGGTCCAGTCGGACTGGA	480
c		V Y T L P P S R D E L T K N Q V S L T C	
•			
		GCCTGGTCAAAGGCTTCTATCCCAGCGACATCGCCGTGGAGTGGGAAGACAATGGGCAGC	540
	481	CGGACCAGTTTCCGAAGATAGGGTCGCTGTAGCGGCACCTCACCCTCTCGTTACCCGTCG	340
С		LVKGFYPSDIAVEWESNGQP	•
		CGGAGAACAACTACAAGACCACGCCTCCCGTGCTGGACTCCGACGGCTCCTTCTTCCTCT	600
	541	GCCTCTTGTTGATGTTCTGGTGCGGAGGGCACGACCTGAGGCTGCCGAGGAAGAAGGAGA	800
С		ENNYKTTPPVLDSDGSFFLY	•
	601	ACAGCAAGCTCACCGTGGACAAGAGCAGGTGGCAGCAGGGGAACGTCTTCTCATGCTCCG	660
	901	TGTCGTTCGAGTGGCACCTGTTCTCGTCCACCGTCGTCCCCTTGCAGAAGAGTACGAGGC	
C		SKLTVDKSRWQQGNVFSCSV	•
	661	TGATGCATGAGGCTCTGCACAACCACTACACGCAGAAGAGCCTCTCCCTGTCTCCGGGTA	720
	901	ACTACGTACTCCGAGACGTGTTGGTGATGTGCGTCTTCTCGGAGAGGGACAGAGGCCCAT	
c	•	MHEALHNHY TQK SLSPGK	•
			•
	721	AAGGTGGAGGTGGTGGAGGTACTTACTCTTGCCACTTCGGCCCGCTGACTTGGGTTT	780
	121	TTCCACCTCCACCACCACCTCCATGAATGAGAACGGTGAAGCCGGCGACTGAACCCAAA	
C		G G G G G G T Y S C H F G P L T W V C	•
		BamHI	
		Bauur	
		GCAAACCGCAGGGTGGTTAATCTCGTGGATCC	
	781	812	
С		CGTTTGGCGTCCCACCAATTAGAGCACCTAGG K P Q G G *	
_			

FIG 14

	X	XbaI		. .		T	•		
	1	TCTAGATTTGTTTTAACTAATTAAAG							
c	_	AGATCTAAACAAAATTGATTAATTTC	CTC	CTTATT				AATGAGAACGG Y S C H	
	61	ACTTCGGCCCGCTGACTTGGGTATGT							
c	01	TGAAGCCGGGCGACTGAACCCATACA F G P L T W V C	ATTC	GGTGTT	CCCC	CACCO	CCTCC	GCCCCCCTGT	•
	121	AAACTCACACATGTCCACCTTGCCCA							
c	121	TTTGAGTGTGTACAGGTGGAACGGGT T H T C P P C P	rcgt	GGACTT	GAGG.	ACCCC	CCTGG	CAGTCAAAAGG	
	101	TCTTCCCCCCAAAACCCAAGGACACC							
c	101	AGAAGGGGGTTTTGGGTTCCTGTGG F P P K P K D T	GGAG	TACTAG	AGGG	CCTGG	GGACT	CCAGTGTACGO	
	244	TGGTGGTGGACGTGAGCCACGAAGAC							
c	241	ACCACCACCTGCACTCGGTGCTTCTC	GGA	CTCCAG	TTCA	AGTTC	ACCAT	GCACCTGCCGC	
		TGGAGGTGCATAATGCCAAGACAAAG							
c	301	ACCTCCACGTATTACGGTTCTGTTTC E V H N A K T K	CGGC	GCCCTC	CTCG	TCATO	TTGTC	GTGCATGGCAG	
		TGGTCAGCGTCCTCACCGTCCTGCAC	CCAG	GACTGG	CTGA	ATGGC	AAGGA	GTACAAGTGC	120
c	361	ACCAGTCGCAGGAGTGGCAGGACGTC V S V L T V L H	GGTC	CTGACC	GACT	TACCO	TTCCT	CATGTTCACG	r
		AGGTCTCCAACAAAGCCCTCCCAGCC	ccc	ATCGAG	AAAA	CCATO	TCCAA	AGCCAAAGGG	2 400
С	421	TCCAGAGGTTGTTTCGGGAGGGTCGC V S N K A L P A	GGGG	TAGCTC	TTTT	GGTAC	AGGTT	TCGGTTTCCC	- 400 -
	401	AGCCCCGAGAACCACAGGTGTACAC							
c	481	TCGGGGCTCTTGGTGTCCACATGTG	GGAC		'AGGG	CCCT	CTCGA	CTGGTTCTTG	
	E 4 1	AGGTCAGCCTGACCTGCCTGGTCAA	AGGC	TTCTAT	CCCA	GCGA	ATCGC	CGTGGAGTGG	3 + 600
c	241	TCCAGTCGGACTGGACGGCCAGTT V S L T C L V K	TCCG	Baagata	GGGT	CGCT	TAGCG	GCACCTCACC	2 -
	601	AGAGCAATGGGCAGCCGGAGAACAA	CTAC	CAAGACC	ACGC	CTCC	GTGCT	GGACTCCGAC	G + 660
С	001	TCTCGTTACCCGTCGGCCTCTTGTTC S N G Q P E N N	GATG	STTCTGG	TGCG	GAGG	CACGA	CCTGAGGCTG	C
	661	GCTCCTTCTTCCTCTACAGCAAGCT	CACC	GTGGAC	AAGA	GCAG	STGGCA	GÇAGGGGAAC	G + 720
С	991	CGAGGAAGAAGGAGATGTCGTTCGA(S F F L Y S K L	GTGG	CACCTG	TTCI	CGTC	CACCGI	CGTCCCCTTG	C
		TCTTCTCATGCTCCGTGATGCATGA	GGC1	CTGCAC	AACC	ACTA	CACGCA	GAAGAGCCTC	T - 780
c	721	AGAAGAGTACGAGGCACTACGTACT F S C S V M H E	CCGA	AGACGTO	TTGG	TGAT	GTGCGI	PCTTCTCGGAG	A
		BamHI							
	700	CCCTGTCTCCGGGTAAATAATGGAT		907					
c	/81	GGGACAGAGGCCCATTTATTACCTA L S P G K +			-				
_									

FIG. 15

	Xì	ai IIO. IO	
	1	TCTAGATTTGAGTTTTAACTTTTAGAAGGAGGAATAAAATATGGGAGGTACTTACT	
b	•	AGATCTAAACTCAAAATTGAAAATCTTCCTCCTTATTTTATACCCTCCATGAATGA	
	61	CCACTTCGGCCCACTGACTTGGGTTTGCAAACCGCAGGGTGGCGGCGGCGGCGGCGGGGGGGG	0
b		GGTGAAGCCGGGTGACTGAACCCAAACGTTTGGCGTCCCACCGCCGCCGCCGCCGCCACCCCGCCGCCGCCGCCAACCGTTTGGCGTCCCACCGCCGCCGCCGCCGCCACCCCCCCC	
	121	TACCTATTCCTGTCATTTTGGCCCGCTGACCTGGGTATGTAAGCCACAAGGGGGTGGGGG + 18	0
b		ATGGATAAGGACAGTAAAACCGGGCGACTGGACCCATACATTCGGTGTTCCCCCACCCCC T Y S C H F G P L T W V C K P Q G G G .	
	181	AGGCGGGGGGACAAAACTCACACATGTCCACCTTGCCCAGCACCTGAACTCCTGGGGGG	0
b		TCCGCCCCCCTGTTTTGAGTGTGTACAGGTGGAACGGGTCGTGGACTTGAGGACCCCCC G G G D K T H T C P P C P A P E L L G G -	
	241	ACCGTCAGTTTTCCTCTTCCCCCCAAAACCCAAGGACACCCTCATGATCTCCCGGACCCC TGGCAGTCAAAAGGAGAAGGGGGGTTTTGGGTTCCTGTGGGAGTACTAGAGGGCCTGGGG TGGCAGTCAAAAGGAGAAGGGGGGTTTTGGGTTCCTGTGGGAGTACTAGAGGGCCTGGGG	0
þ		PSVPLFPPKPKDTLMISRTP-	
	301	TGAGGTCACATGCGTGGTGGACGTGAGCCACGAAGACCCTGAGGTCAAGTTCAACTG+ 36 ACTCCAGTGTACGCACCACCACCACCTGCACTGCGTGCTTCTGGGACTCCAGTTCAAGTTGAC	0
ď		EVTCVVVDVSHEDPEVKFNW-	
	361	GTACGTGGACGGCGTGGAGGTGCATAATGCCAAGACAAAGCCGCGGGAGGAGCAGTACAA CATGCACCTGCCGCACCTCCACGTATTACGGTTCTGTTTCGGCGCCCCTCCTCGTCATGTT	0
þ		Y V D G V E V H N A K T K P R E E Q Y N -	
	421	GTCGTGCATGCCACCAGTCGCAGGAGTGGCAGGACGTGGTCCTGACCGACTTACCGTT	0
b .		S T Y R V V S V L T V L H Q D W L N G K - GGAGTACAAGTGCAAGGTCTCCAACAAAGCCCTCCCAGCCCCCATCGAGAAAACCATCTC	
	481	CCTCATGTTCACGTTCCAGAGGTTGTTTCGGGAGGGTCGGGGGTAGCTCTTTTGGTAGAG	0
b		E Y K C K V S N K A L P A P I E K T I S - CAAAGCCAAAGGCAGCCCCGAGAACCACAGGTGTACACCCTGCCCCCATCCCGGGATGA	
	541	GTTTCGGTTTCCCGTCGGGGCTCTTGGTGTCCACATGTGGGACGGGGGTAGGGCCCTACT KAKGQPREPQVYTLPPSRDE	10
ь		CCTCACCAAGAACCAGGTCAGCCTGACCTGCCTGGTCAAAGGCTTCTATCCCAGCGACAT	. 0
b	601	CGACTGGTTCTTGGTCCAGTCGGACTGGACGACCAGTTTCCGAAGATAGGGTCGCTGTA L T K N Q V S L T C L V K G F Y P S D I	
-		CGCCGTGGAGTGGGAGAGCAATGGGCAGCCGGAGAACAACTACAAGACCACGCCTCCCGT	۰۵
b	661	GCGGCACCTCACCCTCTCGTTACCCGTCGGCCTCTTGTTGATGTTCTGGTGCGGAGGGCA A V E W E S N G Q P E N N Y K T T P P V	
_		GCTGGACTCCGACGGCTCCTTCTTCCTCTACAGCAAGCTCACCGTGGACAAGAGCAGGTG	30
b	721	CGACCTGAGGCTGCCGAGGAAGAAGGAGATGTCGTTCGAGTGGCACCTGTTCTCGTCCAC L D S D G S F F L Y S K L T V D K S R W	. •
	791	GCAGCAGGGGAACGTCTTCTCATGCTCCGTGATGCATGAGGCTCTGCACAACCACTACAC	10
b	701	CGTCGTCCCCTTGCAGAAGAGTACGAGGCACTACGTACTCCGAGACGTGTTGGTGATGTG Q Q G N V F S C S V M H E A L H N H Y T	
		BamHI !	
	841	GCAGAAGAGCCTCTCCCTGTCTCCGGGTAAATAATGGATCC	
þ		Q K S L S L S P G K * SUBSTITUTE SHEET (RULE 26)	

FIG. 16 XbaI TCTAGATTTGTTTTAACTAATTAAAGGAGGAATAACATATGGACAAAACTCACACATGTC AGATCTAAACAAAATTGATTAATTTCCTCCTTATTGTATACCTGTTTTGAGTGTGTACAG C M D K T H T C P -CACCTTGCCCAGCACCTGAACTCCTGGGGGGACCGTCAGTTTTCCTCTTCCCCCCAAAAC 61 -----+ 120 GTGGAACGGGTCGTGGACTTGAGGACCCCCCTGGCAGTCAAAAGGAGAAGGGGGGGTTTTG CPAPELLGGPSVFLFPPRRP-C CCAAGGACACCTCATGATCTCCCGGACCCCTGAGGTCACATGCGTGGTGGTGGACGTGA GGTTCCTGTGGGAGTACTAGAGGGCCTGGGGACTCCAGTGTACGCACCACCACCACCTGCACT K D T L M I S R T P E V T C V V D V S -C GCCACGAAGACCCTGAGGTCAAGTTCAACTGGTACGTGGACGCGTGGAGGTGCATAATG 181 -----+ 240 CGGTGCTTCTGGGACTCCAGTTCAAGTTGACCATGCACCTGCCGCACCTCCACGTATTAC HEDPEVKFNWYVDGVEVHNA-C CCAAGACAAAGCCGCGGGAGGAGCAGTACAACAGCACGTACCGTGTGGTCAGCGTCCTCA GGTTCTGTTTCGGCGCCCTCCTCGTCATGTTGTCGTGCATGGCACACCAGTCGCAGGAGT K T K P R E E Q Y N S T Y R V V S V L c CCGTCCTGCACCAGGACTGGCTGAATGGCAAGGAGTACAAGTGCAAGGTCTCCAACAAAG 301 -----+ 360 GGCAGGACGTGGTCCTGACCGACTTACCGTTCCTCATGTTCACGTTCCAGAGGTTGTTTC V L H Q D W L N G K E Y K C K V S N K A -C CCCTCCCAGCCCCCATCGAGAAAACCATCTCCAAAGCCAAAGGGCAGCCCCGAGAACCAC 361 -----+ 420 GGGAGGGTCGGGGGTAGCTCTTTTGGTAGAGGTTTCGGTTTCCCGTCGGGGCTCTTGGTG L P A P I E K T I S K A K G Q P R E P Q -C AGGTGTACACCCTGCCTCCATCCCGGGATGAGCTGACCAAGAACCAGGTCAGCCTGACCT TCCACATGTGGGACGGAGGTAGGGCCCTACTCGACTGGTTCTTGGTCCAGTCGGACTGGA c V Y T L P P S R D E L T K N Q V S L T C -GCCTGGTCAAAGGCTTCTATCCCAGCGACATCGCCGTGGAGTGGGAGAGCAATGGGCAGC CGGACCAGTTTCCGAAGATAGGGTCGCTGTAGCGGCACCTCACCCTCTCGTTACCCGTCG L V K G F Y P S D I A V E W E S N G Q P -C CGGAGAACAACTACAAGACCACGCCTCCCGTGCTGGACTCCGACGGCTCCTTCTTCCTCT GCCTCTTGTTGATGTTCTGGTGCGGAGGGCACGACCTGAGGCTGCCGAGGAAGAAGGAGA ENNYRTTPPVLDSDGSFFLYc ACAGCAAGCTCACCGTGGACAAGAGCAGGTGGCAGCAGGGGAACGTCTTCTCATGCTCCG TGTCGTTCGAGTGGCACCTGTTCTCGTCCACCGTCGTCCCCTTGCAGAAGAGTACGAGGC c SKLTVDKSRWQQGNVFSCSV-TGATGCATGAGGCTCTGCACAACCACTACACGCAGAAGAGCCTCTCCCTGTCTCCGGGTA ACTACGTACTCCGAGACGTGTTGGTGATGTGCGTCTTCTCGGAGAGGGGACAGAGGCCCAT c M H E A L H N H Y T Q K S L S L S P G K -AAGGTGGAGGTGGCGGAGGTACTTACTCTTGCCACTTCGGCCCACTGACTTGGGTTT TTCCACCTCCACCACCGCCTCCATGAATGAGAACGGTGAAGCCGGGTGACTGAACCCAAA G G G G G G T Y S C H F G P L T C GCAAACCGCAGGGTGGCGGCGGCGGCGGCGGTGGTACCTATTCCTGTCATTTTGGCCCGC CGTTTGGCGTCCCACCGCCGCCGCCGCCACCATGGATAAGGACAGTAAAACCGGGCG C KPQGGGGGGGTYSCHFGPL-BamHI -TGACCTGGGTATGTAAGCCACAAGGGGGTTAATCTCGAGGATCC 841 ----- 884 ACTGGACCCATACATTCGGTGTTCCCCCAATTAGAGCTCCTAGG T W V C K P Q G G .

FIG. 17A

[AatII sticky end] (position #4358 in pAMG21)

- 5' GCGTAACGTATGCATGGTCTCC3' TGCACGCATTGCATACGTACCAGAGG-
- -CCATGCGAGAGTAGGGAACTGCCAGGCATCAAATAAAACGAAAGGCTCAGTCGAAAGACT--GGTACGCTCTCATCCCTTGACGGTCCGTAGTTTATTTTGCTTTCCGAGTCAGCTTTCTGA-
- -GGGCCTTTCGTTTTATCTGTTGTTGTCGGTGAACGCTCTCCTGAGTAGGACAAATCCGC--CCCGGAAAGCAAAATAGACAACAAACAGCCACTTGCGAGAGGACTCATCCTGTTTAGGCG-
- CGGGAGCGGATTTGAACGTTGCGAAGCAACGCCCGGAGGGTGGCGGGCAGGACGCCCGC GCCCTCGCCTAAACTTGCAACGCTTCGTTGCCGGGCCTCCCACCGCCCGTCCTGCGGGCG -
- CATANACTGCCAGGCATCANATTANGCAGNAGGCCATCCTGACGGATGGCCTTTTTGCGT GTATTTGACGGTCCGTNGTTTNATTCGTCTTCCGGTNGGACTGCCTNCCGGANANCGCN -

AatII

- TTCTACAAACTCTTTTGTTTATTTTTCTAAATACATTCAAATATGGACGTCGTACTTAAC AAGATGTTTGAGAAAAAAAAAAAAAAAAAGATTTATGTAAGTTTATACCTGCAGCATGAATTG -
- TTTTAAAGTATGGGCAATCAATTGCTCCTGTTAAAATTGCTTTAGAAATACTTTGGCAGC AAAATTTCATACCGTTAGTTAACGAGGACAATTTTAACGAAATCTTTATGAAACCGTCG -
- GGTTTGTTGTATTGAGTTTCATTTGCGCATTGGTTAAATGGAAAGTGACCGTGCGCTTAC CCAAACAACATAACTCAAAGTAAACGCGTAACCAATTTACCTTTCACTGGCACGCGAATG -
- -TACAGCCTAATATTTTTGAAATATCCCAAGAGCTTTTTCCTTCGCATGCCCACGCTAAAC-ATGTCGGATTATAAAAACTTTATAGGGTTCTCGAAAAAGGAAGCGTACGGGTGCGATTTG-
- -GATAATTATCAACTAGAGAAGGAACAATTAATGGTATGTTCATACACGCATGTAAAAATA --CTATTAATAGTTGATCTCTTCCTTGTTAATTACCATACAAGTATGTGCGTACATTTTTAT -
- AACTATCTATATAGTTGTCTTTCTCTGAATGTGCAAAACTAAGCATTCCGAAGCCATTAT TTGATAGATATATCAACAGAAAGAGACTTACACGTTTTGATTCGTAAGGCTTCGGTAATA -
- TAGCAGTATGAATAGGGAAACTAAACCCAGTGATAAGACCTGATGATTTCGCTTCTTTAA ATCGTCATACTTATCCCTTTGATTTGGGTCACTATTCTGGACTACTAAAGCGAAGAAATT -
- TTACATTTGGAGATTTTTTATTTACAGCATTGTTTTCAAATATATTCCAATTAATCGGTG AATGTAAACCTCTAAAAAATAAATGTCGTAACAAAAGTTTATATAAGGTTAATTAGCCAC -
- AATGATTGGAGTTAGAATAATCTACTATAGGATCATATTTTATTAAATTAGCGTCATCAT TTACTAACCTCAATCTTATTAGATGATATCCTAGTATAAAATAATTTAATCGCAGTAGTA -
- AATATTGCCTCCATTTTTTAGGGTAATTATCCAGAATTGAAATATCAGATTTAACCATAG TTATAACGGAGGTAAAAAATCCCATTAATAGGTCTTAACTTTATAGTCTAAATTGGTATC -
- AATGAGGATAAATGATCGCGAGTAAATAATATTCACAATGTACCATTTTAGTCATATCAG-- TTACTCCTATTTACTAGCGCTCATTTATTATAAGTGTTACATGGTAAAATCAGTATAGTC -

- GCAAGTTTTGCGTGTTATATATATCATTAAAACGGTAATAGATTGACATTTGATTCTAATAA CGTTCAAAACGCACAATATATAGTAATTTTGCCATTATCTAACTGTAAACTAAGATTATT -

FIG. 17B

- ATTGGATTTTTGTCACACTATTATATCGCTTGAAATACAATTGTTTAACATAAGTACCTG-
- -TAACCTAAAAACAGTGTGATAATATAGCGAACTTTATGTTAACAAATTGTATTCATGGAC -
- $\hbox{-} TAGGATCGTACAGGTTTACGCAAGAAAATGGTTTGTTATAGTCGATTAATCGATTTGATT-\\$
- -ATCCTAGCATGTCCAAATGCGTTCTTTTACCAAACAATATCAGCTAATTAGCTAAACTAA
- -CTAGATTTGTTTTAACTAATTAAAGGAGGAATAACATATGGTTAACGCGTTGGAATTCGA-
- -GATCTAAACAAAATTGATTAATTTCCTCCTTATTGTATACCAATTGCGCAACCTTAAGCT-
- Sacii -GCTCACTAGTGTCGACCTGCAGGGTACCATGGAAGCTTACTCGAGGATCCGCGGAAAGAA-
- -CGAGTGATCACAGCTGGACGTCCCATGGTACCTTCGAATGAGCTCCTAGGCGCCTTTCTT
- GAAGAAGAAGAAGCCCGAAAGGAAGCTGAGTTGGCTGCCACCGCTGAGCAATA CTTCTTCTTCTTTCTGGGCTTTCCTTCGACTCAACCGACGACGACGGCGACTCGTTAT -
- -ACTAGCATAACCCCTTGGGGCCTCTAAACGGGTCTTGAGGGGTTTTTTGCTGAAAGGAGG-
- -TGATCGTATTGGGGAACCCCGGAGATTTGCCCAGAACTCCCCAAAAAACGACTTTCCTCC-
- -AACCGCTCTTCACGCTCTTCACGC 3!
- [SacII sticky end]
- -TTGGCGAGAAGTGCGAGAAGTG 5'
- (position #5904 in pAMG21)

FIG.18A - 1

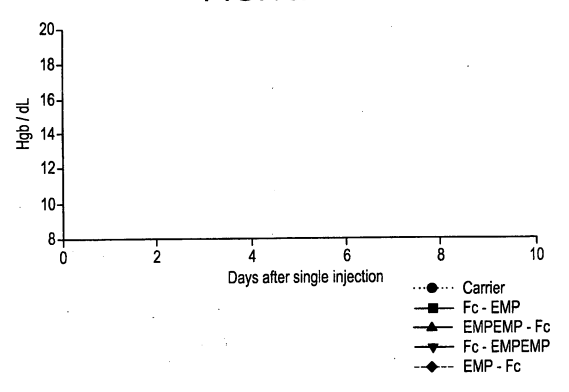


FIG.18A - 2

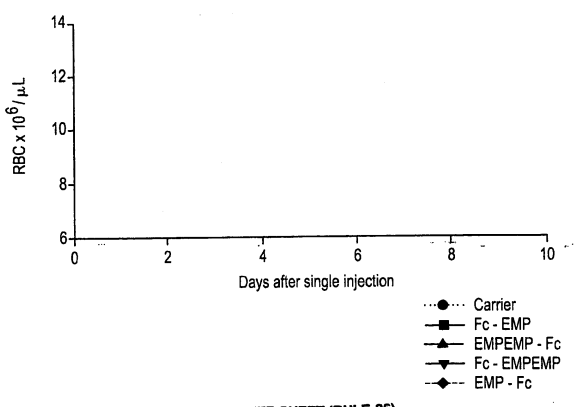


FIG.18A - 3

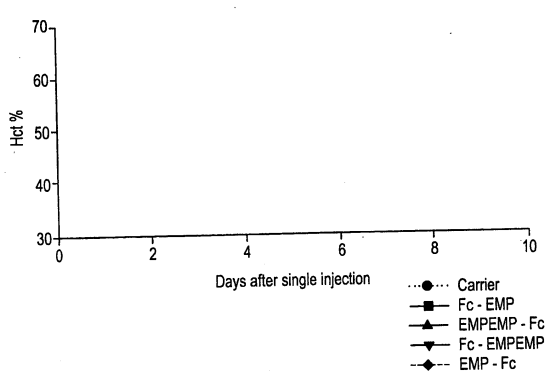


FIG.18B - 1

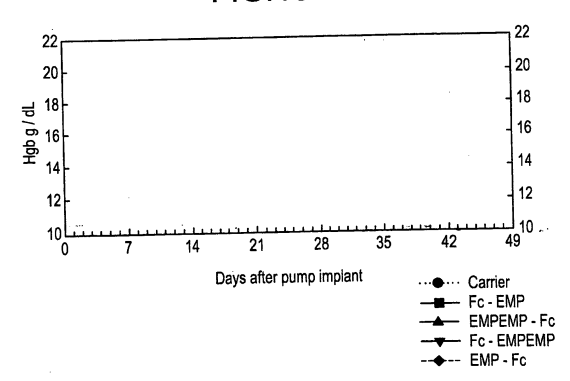


FIG.18B - 2

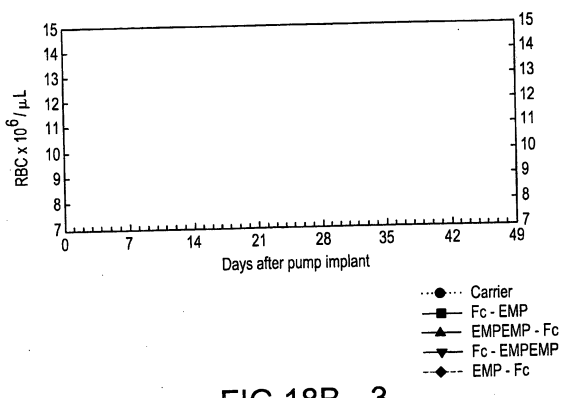
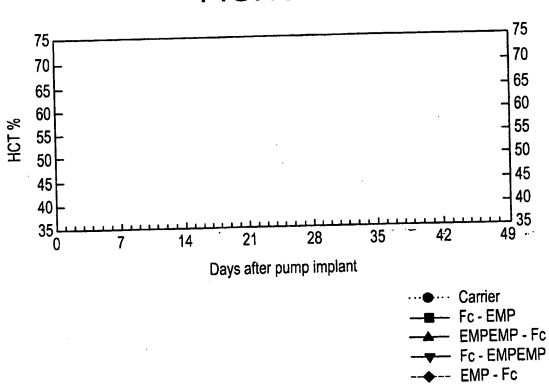


FIG.18B - 3



	NdeI								F	-[(3		19)Δ	\							
	1							-+-			+	• • •			+						CCG + CGC	60
			м	D	K				С								L			_	P	•
	61										+				- +							120
		AG:					igge P							L	M	I	S	R	T	P	CTC E	
1		S												_			CAAC	TTC	AAC	TGC	TAC	180
	121							+ -			+				- + -						CATG	100
1		•	-	С	-	V		D	V	S	Н	_	ď·	P	E	V	K	F	N	W	Y	•
	181							+ -			4				- + -						CAGC + GTCG	240
3.		V	D	G	v	E	V	Н	N	A	ĸ	T	K	P	R	E	E	Q	Y	N	S	-
	241				- 4 -							+	• • •		-+-			• • •				300
		TG	CAT	GGC						_			H	0	D	w	L	N.	G	ĸ	CCTC E	-
a		T	Y	R	V	v 	S 	v 	L	T	V	L ccc		-	_		_		_		_	
	301											+			• + •	. 					CAAA GTTT	200
á		Y	ĸ	С	K	V	s	N	ĸ	A	L	P	A	P	I	E	K	-	I	•	K	•
	36:						-					+- •			• • •						GCTG CGAC	320
a		A	ĸ	G	Q	P	R									P			D	E	L	•
_	42																				CGC(
																					A	
a								ma	יייי א	ccc	car	CA	1 01	ACT!	ACA	AGA	CAC	GCC	TC	CG:	rgcT(3
	48	1 - C	ACC	TCA	CCC	TCT	CGT	rac	CGI	CGC	CC	rct'	TGT'	TGA'	TGT	TCT	GGT	3CGC	BAG	GGC/	ACGA	
а		v	E	W	E	s	N	G	Q	P	E	N	N	Y	K	T	T	P	P	V	L	^ · -
		_										. + -			• • •				•		GGCA	
	54	C	TGA	GGC	TGC	CGA	GGA	AGA	AGG	AGA'	rgt(CGT	TCG	AGT	GGC	ACC	IGI	1010	-G1,	CCA	CCGI	•
		_					2	F	τ,	Y	S	ĸ	L	т	. v	r D	K	S	R	W	Q	•

FIG. 19B

	Q	G	N	v	F	S	С	S	V	M	H	E	A	L	Н	N	Н	Y	T	Q	
661							+				+••			-+-			+			CTAC	•
											ACC G									'GATG Y	
	K	3	u	3	_		•	Ĭ	•		mHI				•						

FIG. 20A

		INCL																				
		CAT	ATG	GAC	TTC	CTC	CC	GCA	CTA	CAAA	AAC	ACC	TCT	CTG	GGT	CAC	CGT	'CCC	GGT	rgga	GGC	60
											4				+						CCG	00
_									Y								R	P	G	G	G	
a					1		nc a	CAC	አጥርና	ጥሮሮን	אריכי אריכי	ኮጥርር	CCA	.GCA	CCT	'GAA	CTC	CTC	GGG	3GG∤	ACCG	
															. +			· - T ·			•	120
		CCA	CCC	CTC	GTT'	rtg.	AGT	GTG	TAC	AGG'	rgg	AAC	انافاذ	rcG1	rGGA						rggc	
a		_	_		ĸ					P	P	С	P	A	P	E	L	L	G	G	P	•
	121	TCA	GTI	TT	CCT	СТТ	ccc	ccc	AAA:	ACC	CAA	GGA	CAC	CCT	CATO	ATC	TC	CCG	GAC	CCC'	TGAG	180
	121	AGT	CA	LAA!	-+- GGA	GAA	GGG	GGC	TTT	тgg	GTT	CCT	GTG	GGA	GTAC	CTAC	SAG	GGC	CTG	GGG.	ACTC	
a		s	v	F					K			D		L		I	_	R	T	P	_	•
-		GT/C	יארי	ነጥር	CGT	GGT	GGT	rgg!	ACGI	GAG	CCA	CGA	AGA	CCC'	TGA	GT(CAA	GTT	CAA	CTG	GTAC	240
	181																				CATG	
		CAG	STG:	rac	GCA	CCA		3CC.		_					E						Y	•
a		V	T	C	V	V	٧	Þ	٧	_	Н	_	_	_						•••	-	
		GT	GGA	CGG	CGI	'GG?	\GG'	rgc	ATA	ATGC	CAA	GAC	AAA	GCC	GCG	GGA	GGA	.GCA	GT?	CAA	CAGC	300
	241	CAC	 CCT	GCC	· + · GCA	ACC1	rcc	ACG	TAT:	raco	GTI	CTC	TTT	'CGG	CGC	CCT	CCI	CGI	'CA'	rgti	GTC	}
•		v	'n	G	v	E	v			. д		T		P						N		-
a		•	_			ncci	የረግ አ	ccc	ጥ ()	TCAC	ccgi	rcci	rgc#	CCA	\GGA	CTG	GC1	'GA	\TG(GCA!	AGGA	3
	301																					
		TG	CAT	'GG(CAC	ACC	AGT														rcct(•
a		T	Y	R	-		_	-	L						D					K		_
																					CCAA	
	361	AT	GTI	CA	+ CGT'	TCC	AGA	GGI	TGT	TTC	GGG.	AGG	GTC	GGG	GGT/	AGCT	CT'	rtt	GGT.	AGA	GGTT'	T
a			к						ı K							E	K	_	_	_		-
-		GC.	ו מיזי	A A C	ccc	AGC	ccc	GAC	AAC	CAC	AGG	TGT	ACA	CCC'	TGC	ccc	CAT	ccc	GGG	ATG	AGCT	G 400
	423																				TCGA	
а																					L	
		A(CA	AGA	ACC	AGG	STC	AGC	CTG	ACCI	GCC	TGG	TCA	AAG	GCT'	TCT.	ATC	CCA	GCG +	ACA	TCGC	±. 540
	48	1 - ·	 GGT'	TCI	TGC	TCC	'AG'	TCG	GAC:	rgg?	CGG	ACC	AGT	TTC	CGA	AGA	TAG	GGT	CGC	TGT	AGC	G.
a		т	ĸ	N	1 ()	,	S	L S	r	1	, ,	7 K	; G	F	Y	F	2	S I) 1	A	•
•										~				3.00	ארא	AGA	CCZ	CGC	CTC	ccc	TGC	ľG
	54	1 -			• • •				-+-	cmc/		- + ·	ייטיטין יייטיטיין	 РФС 2	+ \ጥር፡ጥ	TCT	'GG'	rgc	· + · · 3GA(GGG	CACG	+ 600 AC
а		٧	E	: 1	v 1	E	S	N	G	Q :	P !	Ε]	1 1	1 ,	e K	. 1			5	-	V L	

FIG. 20B

601				-+-			+				+			-+-			+			CGTC	660
	D	s	D	G	s	F	F	L	Y	S	K	L	т	V	D	K	S	R	W	Q	•
	_	•	i		_					. .						~~ .					
	CA	GGG	GAA	CGI	CTI	CTC	ATG	CTC	CGT	'GA'I	GCA	TGA	GGC	TCT	GCA	CAA	CCA	CTA	CAC	GCAG	720
661	GT	CCC	CTI	GCA	GAA													GAT	GTC	CGTC	
-	-																		т	0	
	Q	G	N	V	F	S	С	S	٧	M	H	E	A	ы	п	IA	п	1	1	V	
										n.	ımH I										
											-										
	AA	GAC	CCI	CTC	CCI	CTC	TCC	:GG(TAP	\AT?	ATC	GAT	rccc	CGG	} · 76	: 1					
721		·	cai	· - + ·		·	+ 2AGC	CCC	ירב. וידא:	'TA'	TAC	CT	\GG(GCC		, 1					
-	11	CIC	,GGr	JOAC	3002	CARC	,,,,,,														
	v	c	τ.	Q	t.	g	P	G	K	*			•								

FIG. 21A

		eI 									com	men	CCN	CCT	ecco	C A A	crec	ירייר	GGG	CCA	cce	
	1				. +			+ -			+				+			• • • •	CCC		+	60
		GT	ATAC	CTC	3'I''I''I	l"I'G#	AGTC														P	_
			M	_	K	T	Н	Ť	_	_	P	_				_	_	L		G	-	•
	61				- 4		 .	+ -			+				.+			+			GAG	120
	0.1	AG'	rca(SAA:	GGA(GAAC	GGG	GGG	rtti	`GGG	TTC	CTC	TGG	GAG	TAC	TAC	GAG	3GC(CTGG	GG#	CTC	
Ļ		s	v	F	L	F	P	P	ĸ	P	K	D	T	L	M	I	S	R	T	P	E	•
		GT	CAC	ATG	CGT	GGT	GGT	GGA	CGTC	AGC	CAC	GA/	GAC	CCI	rgac	GT	CAA	3TT(+	CAAC	TGC	TAC	180
	121	CA	GTG'	TAC	GCA	CCA	CCA	CCT	GCAC	TCC	GTC	CTI	CTC	GGZ	CTC	CAC	GTT(CAA	GTT(BACC	CATG	
l		v	т	С	v	v	v	D	v	s	H	E	D	P	E	v	K	F	N	W	Y	•
		GT:	GGA	CGG	CGT	GGA	GGT	GCA'	TAAT	rgco	CAAC	BAC	\AA(GCC(GCG(GA (GGA(GCA	GTA	CAAC	CAGC	240
	181							+						• • • •	-+-		• • •	•••			GTCG	240
									N	_	ĸ	T	ĸ	P	R	E	E	0	Y	N	s	-
1		-	_		v							_					ርርጥ ርርጥ	CAA	ጥርር	CAA	GGAG	
	241							+				+		• • •	-+-		• • •				GGAG	300
		TG	CAT	GGC	ACA	CCA	GTC	GCA	GGA	GTG					_						CCTC	
a		T	Y		V	V	S	-	L	T		L		**	D	•••	L	N	G	K	E	-
	301							+				+			-+-			7				360
	201	ΑΊ	GTI	CAC	GTT	CCA	GAG	GTI	GTT	TCG	GGA	GGG	TCG	GGG	GTA	GCT	CTI	'TTC	GTA	GAG	GTTT	
a		Y	к	С	K	v	S	N	ĸ	A	L	P	A	P	I	E	K	T	I	S	K	-
		GC	CA	AGG	GC#	AGCC	ccc	AG!	ACC	ACA	.GGT	GTA	CAC	CCT	GCC	ccc	ATC	ccc	GGA	TGA	GCTG	420
	361	CC	GT1	rtco	- + - CG7	rcgo	GGG	CTC	rtgg	TGT	CCA	CAT	GTG	GGA	CGG	GGG	TAC	GGG	CCI	ACT	CGAC	
a		Α	к	G	Q	P	R	E	P	Q	V.	Y	T	L	P	P	s	R	D	E	L	•
-		A	CA	AGA/	ACC/	AGGT	rcac	3CC!	rgac	CTC	CCI	'GG'I	CAA	AGG	CTI	CT	ATC	CAC	GCG#	CAT	CGCC	400
	421																				AGCGG	
																					A	
а																					rgcto	
	481											• + - •		• • • •	+	• • •					7	,, 340
																					ACGAC	
a																					L	
	E A .											-+-	* .		+				-		GGCAC	000
	34.	C	TGA	.GGC	TGC	CGA	GGA	AGA	AGG	AGA'	rgt	CGT'	rcg	AGT	GGC.	ACC'	TGT	TCT	ÇGT	CCA	CCGTC	•
a		D	S	ם	G	s	F	F	L	Y	S	K	L	T	V	D	K	S	R	W	Q	•

FIG. 21B

	GT	CCC	CTT	GCA	JAA	JAG														CGTC
	Q	G	N	V	F	S	С	S	V	M	Н	E	A	L	Н	N	Н	Y	T	Q
661																				GGGT GCCCA
			L				P				G						W	T	P	G
721							4			rgT/	I Hme TAA - + -	GAT	-			763	3			

FIG. 22A

		Nde	eΙ																		
		CAT	ATGT	TCGA	LATG			GGG							CTC	CCC	CTC	GG.	rgg,		60
	1	GTA:	TACA	AGCI	TAC						•				AGAC	CGGC	CGA	CCZ	ACCI		60
a		1	M F	E	W	T	P	G	Y	W	Q	P	Y	A	L	P	L	G	G	G	•
			GGGG	ACAA	AAC	TCA		ATG										GG(GG/		120
	61	CCA	cccc	TGTI	TTG	AGT												CCC	ccc		120
a		G	G D	к	T	н	T	С	P	P	С	P	A	P	E	L	Ļ	G	G	P	•
			GTTT		CTI										GAT	CTC	CCG	GAC	ccc		180
•	121	AGT	CAAA												CTA	GAG	GGC	CTG	GGG		100
a		s ·	v f	L	F	P	P	ĸ	P	ĸ	D	T	L	M	I	S	R	T	P	E	-
		GTC.	ACAT	GCGI	rggī	GGT	GGA	CGT	GAG	CCA	CGA	AGA	CCC	TGA	GGT(CAA	GTT(CAA	CTG	GTAC	240
	181	CAG	TGTA	CGC	ACCA	CCA	CCI	GCA	CTC	GGT	GCT	TCT	GGG	ACT	CCA	GTT(CAA	GTT	GAC	CATG	240
a		v	тс	. v	v	v	D	v	s	H	E	D	P	E	V	ĸ	F	N	W	Y	•
		GTG	GACG	GCG:	rgg <i>i</i>	\GG1	'GCA	\TAA	TGC	CAA	GAC	AAA	.GCC	GCG	GGA(GGA	GCA	GTA	CAA	CAGC	200
	241	CAC	CTGC	CGC2	ACCI	rcc?	CGI	TTA	ACG	GTT	+ CTG	TTT	CGG	CGC	CCT	CCT	CGT	CAT	GTT	GTCG	300
a		v	D G	v	E	v	Н	N	A	K	T	ĸ	P	R	E	E	Q	Y	N	s	•
		ACG	TACC	GTG	rggi	CAC	GCG1	rcci	CAC	CGT	CCT	GCA	CCA	GGA	CTG	GCT	GAA	TGG	CAA	GGAG	260
	301	TGC	ATG	CAC	ACC!	AGTO	CGC	AGGA	GTG	GCA	GGA	CGT	GGT	CCT	GAC	CGA	CTT	ACC	GTT	CCTC	360
a		T	Y F	v ک	v	S	v	L	T	V	L	н	Q	D	W	L	N	G	K	E	-
		TAC	AAGI	rgca.	AGG'	rcto	CA	ACAA	AGC	CCI	ccc	AGC	ccc	CAT	CGA	GAA	AAC	CAT	CTC	CAAA	420
	361	ATG	TTC	CGT'	TCC	AGA	GT'	rgti	TCC	GG?	\GGG	TCG	GGG	GTA	GCT	CTT	ТТG	GTA	GAG	GTTT	420
a		Y	K C	: к	v	S	N	K	A	L	P	A	P	I	E	K	T	I	S	K	-
		GCC	AAA	GGC.	AGC	ccc	GAG/	AAC	CAC	\GG1	GTA	CAC	CCI	GCC	CCC	ATC	CCG	GGA	TGA	GCTG	480
	421	CGG	TTT	CCG	TCG	GGG	CTC'	rtgo	TG	rcc.	CAT	GTG	GGA	CGG	GGG	TAG	GGC	CCT	ACT	CGAC	400
a		A	K (3 Q	P	R	E	P	Q	v	Y	T	L	P	P	S	R	D	E	L	•
		ACC	AAG	AACC	AGG'	TCA	GCC'	TGA	CTC	GCC1	rggi	CAA	AAGO	CTI	'CTA	TCC	CAG	CGA	CAT	CGCC	E 4 0
	481	ТĞС	TTC	+ TTGG	TCC	AGT(CGG	act(GA(CGG2	ACC!	GTI	rTC	GAA	GAT	AGG	GTC	GCI	GTA	.GCGG	540
a		т	K I	. Q	v	S	L	T	С	L	V	K	G	F	Y	P	s	D	I	A	-
		GTC	igagʻ	rggg	AGA	GCA.	ATG	GGC	AGC	CGG	AGA	ACA/	ACT	ACA?	GAC	CAC	:GCC	TCC	CGI	GCTG	
	541	CAC	CTC	ACCC	TCT	cgt	TAC	+ CCGʻ	rcg	GCC'	rct:	rgti	rga:	rgT1	CTC	GTO	CGC	AGC	GCA	CGAC	600
а		v	E 1	W E	: S	N	G	Q	P	E	N	N	Y	ĸ	T	т	P	P	v	L	

FIG. 22B

	601				-+-			+				+			-+-			+			CGTC	660
a		D	s	D	G	s	F	F	L	Y	S	K	L	T	V	D	K	s	R	W	Q	•
	661		<i>-</i>		-+-			+				+			-+-			+	·		GCAG GCGTC	720
a		Q	G	N	v	F	s	С	s	V	M	Н	E	A	L	Н	N	Н	Y	T	Q	•
	721				CTC			+			ATA	+	GAT		757	7						

FIG. 23A

	Nd	leI																				
	1				-+-		 .	+ -	· ·	·	· ·	+ ·	· · · ·	- - -	+			-+-				60
		GTA				rtg/												L.			GGC	
3			M	D	K	T	н	т	С	P	₽	•	P	A	P	E	L	_	G	G	P	•
	61				-+-			+	• - • ·		• • •	+			+			-+-			GAG + CTC	120
a		S	V	F	L	F	P	P	K	P	K	D	T	L	M	I	S	R	T	P	Ē.	•
																					TAC	
	121																				ATG	180
a	•	v	T	С	v	v	v	D	v	s	н	E	D	P	E	v	ĸ	F	N	W	Υ .	
		GTO	GAG	CGG	CGT	GGA(GGT	GCA!	raa:	rgc(CAA	GAC	AAA(3CC(GCGC	3GA(GAG	CAC	TA(CAAC	CAGC	
	181					CCT													CAT	TTC	TCG	240
a		v	ם	G	v			н				т								N	S	
_				_	· •	_	•		-	73.0	~~~	CCTI	: ::::::::::::::::::::::::::::::::::::		7C N C	~mc/	2CTC	- יא אי	raaa	7 A A C	GAG	
	241				-+-			+				+			-+-			+ -	• • •		CTC	300
															D		L			K	E	
a		T	Y 		v	•	_	V		-	-	L		•	_		_		_		_	
	301				-+-			+				+			•+•			+				360
		ATO	STT(CAC	GTT	CCA	GAG	GTT(GTT'	rcg	GGA	GGG'	rcg(·	GGG(GTA(GCT(CTT'	rtg(3TA(3AG(TTT	
a		Y	K	С	K	V	S	N	K	Α	L	P	A	P	I	E	K	T	I	S	K	•
	361		:AA	AGG	GCA	GCC	CCG.					GTA			GCC(-+-	CCC	ATC	CCG(GA'	rgac	CTG	420
		CGC	3TT	TCC	CGT	CGG	GGC	TCT"	TGGʻ	TGT	CCA	CAT	GTG	GGA	CGG	GGG	rag	3GC(CT	ACT	CGAC	
a		A	K	G	Q	P	R	E	P	Q	V	Y	T	L	P	P	S	R	D	E	L	•
		ACC	CAA	GAA	CCA	GGT	CAG	CCT	GAC	CTG	CCT	GGT	CAA	AGG	CTT	CTA!	rcc	CAG	CGA	CAT	CGCC	480
	421	TG	GTT	CTT	ggt	CCA	GTC	GGA	CTG	GAC	GGA	CCA	GTT	TCC	GAA	GAT	AGG	GTC	GCT	GTA(GCGG	400
a		T	ĸ	N	Q	v	s	L	T	С	L	V	ĸ	G	F	Y	P	s	D	I	A	•
		GT	GGA	GTG	GGA	GAG	CAA	TGG	GCA	GCC	GGA	GAA	CAA	CTA	CAA	GAC	CAC	GCC'	TCC	CGT	GCTG	E 4.0
	481	CA	CCT	CAC	CCT	CTC	GTT	ACC	CGT	CGG	CCT	CTT	GTT	GAT	GTT	CTG	GTG	CGG.	ĀGG	GCA	CGAC	340
a		v	E	W	E	S	N	G	Q	P	E	N	N	Y	K	T	T	P	P	v	L	-
		GA	ርጥሮ	CGA	റദേ	CTC	стт	CTT	CCT	СТА	CAG	CAA	GCT	CAC	CGT	GGA	CAA	GAG	CAG	GTG	GCAG	
	541				-+-			+				+			•+-			+			CGTC	600
a																					Q	-
~		_	_	_	-		-	-	_	-	-		-								-	

FIG. 23B

	601	CA	GGG	GAA	CGT	CTT	CTC	ATG	CTC	CGT											+	660
	001	GT	CCC	CTT	GCA	GAA	GAG	TAC	GAG	GCA	CTA	CGT	ACT	CCG	AGA	CGT	GTT	GGT	'GAI	GTG	CGTC	
a		Q	G	N	v	F	S	С	S	V	M	Н	E	A	L	Н	N	Н	Y	T	Q	•
•	661				•+•			+				+			-+-			+			TGAC + ACTG	720
a		ĸ	s	L	S	L	s	P	G	K	G	G	G	G	G	V	E	P	N	С	D	-
																_	amH	ī		_		
	721				-+-			+			TTT	+			-+-			• • •		• 77	73	
a		I	н	v	M	W	E	W	E	С	F	E	R	L	*							

FIG. 24A

	NC	ieI																				
	1	CA:	TAT(GGT'	TGA.	ACC	GAA	CTG	TGA	CAT		rgt:				ATG	GA/	TGI	rTT.	'GAA	CGT	60
		GT	ATA	CCA	ACT'	TGG	CTT	GAC.	ACT	GTA(GT.	ACAZ	ATAC	CAC	CT	raco	CTT	ACA	LAA.	CTI	'GCA	
a			M	V	E	P	N	C	D	I	Н	V	M	W	E	W	E	С	F	E	R	•
	61			TGGʻ													_				CTC	120
	0.1																				GAG	120
a		L	G	G	G	G	G	D	K	T	H	T	С	P	P	С	P	A	P	E	L	-
	121	CT																			TCC	180
	121	GA																			BAGG	100
a		L	G	G	P	S	V	F	L	F	P	P	K	P	K	D	T	L	M	I	s	•
																					AAG	242
	181																				TTC	240
a		R	т	P	E	v	T	С	v	v	v	D	v	s	н	E	D .	P	È	v	K	-
			CAA	CTG																	GAG	
	241		GTT(GAC																	CTC	300
a		F	N	W	Y	v	D	G	v	E	V	н	N	A	ĸ	T	ĸ	P	R	E	E	•
	301	••			-+-			+				+			-+-			· - + ·	· ·	· ·	GCTG	360
		GT		GTT(ACA(_				_	_		CGAC	
a		Q	Y	N	S	T	Y	R	V	V	S	V	L	T	V	L	H	Q	D	W	L	•
	361				-+-			+				+			-+-			+ -			GAAA +	420
		TT.	ACC	GTT	CCT	CAT	GTT	CAC	GTT	CCA	GAG	GTT(GTT'	rcg	GGA(GGG'	rcg	3GG(STA(3CT(CTTT	
a		N	G	K	E	Y	K	С	K	V	S	N	K	A	L	P	A	P	I	E	K	•
	421				-+-	• • •	•	+	· · ·	·		+••			-+-		• • •	+				480
		TG	GTA	GAG	GTT	TCG	GTT	TCC	CGT	CGG	GGC	TCT	TGG'	rg T	CCA	CAT	GTG(GGA(CGG	GGG'	FAGG	
a		T	I	S	K	A	K	G	Q	P	R	E	P	Q	V	Y	T	L	P	P	S	•
	481								CCA											CTA!	rccc +	340
		GC	CCT	ACT	CGA	CTG	GTT	CTT	GGT	CCA	GTC	GGA	CTG	GAC	GGA	CCA	GTT'	rcc	GAA(GAT	AGGG	
a		R	D	E	L	T	K	N	Q	V	S	L	T	С	L	V	K	G	F	Y	P	-
	541	AG	CGA	CAT	CGC	CGT	GGA	GTG	GGA	GAG	CAA	TGG +	GCA	GCC	GGA	GAA	CAA	CTA(CAA	GAC	CACG	600
																					GTGC	
a		S	D	I	A				E							N	N	Y	K	T	T	-
						~																

FIG. 24B

	601	••			-+-			+	·		· · ·	+			•+-			+			GTTC	660
a		P	P	V	L	D	· S	D	G	S	F	F	L	Y	S	K	·L	T	V	D	K	-
	661				-+-			+	· • • •		• • •	+			-+-	• • •		+			CAAC GTTG	720
a		s	R	W	Q	Q	G	N	V	F	S	С	S	V	M	Н	E	A	L	Н	N	•
	721				-+-			+				+			-+-	ACI		II \GGA 	· • • •	77	3	
		u	v	4	0	ĸ	S	T.	S	τ.	S	P	G	K	*							

FIG. 25A

	No	leI																			
	1	CATA	TGGA																		60
	_	GTAT																			00
a		М	D	ĸ	T	Н	T	c	P	P	С	P	A	P	E	L	L	G	G	P	-
	£ 1	TCAG	тстт																		120
	01	AGTC																			120
a		s v	F	L	F	P	P	ĸ	P	ĸ	D	T	L	M	I	s	R	T	P	E	-
	1 2 1	GTCA																			180
	121	CAGT																			180
a		v r	С	v	V	v	D	V	S	Н	E	Ď	P	E	v	K	F	N	W	Y	•
		GTGG																TAC	CAAC	AGC	040
	181	CACC																ATC	TTC	TCG	240
a		v D	G	v	E	v	н	N	A	ĸ	T	ĸ	P	R	E	E	Q	Y	N	S	-
		ACGT																			
	241	TGCA																			300
a		T Y	R	v	v	s	v	L	T	v	L	H	Q	ס -	W	L	N	G	ĸ	E	-
		TACA																			
	301	ATGT																			360
a		у к	C	K	V	s	N	ĸ	A	L	P	A	P	I	E	K	T	I	s	ĸ	•
		GCCA																			400
	361	CGGT																			420
a		A K	G	Q	P	R	E	P	Q	V	Y	T	L	P	P	S	R	D	E	L	-
		ACCA																			400
	421	TGGT	TCTI																		480
a		т к	N	Q	v	S	L	T	С	L	V	K	G	F	Y	P	s	D	I	A	•
		GTGG																			
	481	CACC																			540
a		V E	W	E	s	N	G	Q	P	E	N	N	Y	K	T	T	p	P	v	L	-
		GACT	CCGA	CGG	CTC	CTT	CTT	ССТ	CTA	CAG	CAA	GCT	CAC	CGT	GGA(CAAC	GAG(CAG	3TG(GCAG	
	541	CTGA	GGCI	GCC	GAG	GAA	GAA	GGA	GAT	GTC	+ GTT	CGA	GTG(GCA	CCT	STT(CTC	GTC	CAC	CGTC	600
a		D S	D	G	s	F	F	L	Y	S	к	L	T	v	D	ĸ	s	R	W	Q	-

FIG. 25B

	GT	CCC	CTT																	CGTC
	Q	G	N	V	F	S	С	S	V	M	H	E	A	L	Н	N	H	Y	T	Q
661				-+-			+				+			-+-			+			GGGT CCCA
	K	S	L	s	L	3	P	G	K	G	G	G	G	G	С	T	T	H	W	G
						I Hm.														
721			CCT	-+-			4			·	748)								

FIG. 26A

	No	ieI																				
	1				-+-			+			CACC	 -			+		• • •	-+-			+	60
		GT	ATA	CAC	GTG	GTG	GGT	GAC	CCC	AAA(GTGC	3GA(CAC	3CC?	ACCI	CCG	CCA	CCC	CTG	TTT	CCA	
1			M	С	T	T	Н	W	G	F	T	L	С	G	G	G	G	G	D	K	G	-
	61		- <u>-</u> -		-+-			+				 -			-+			-+-		• - •	+	120
		CC	rcc	GCC	ACC	CCT	GTT	TTG.	AGT	3TG	TAC	AGG'	rgg	AACC	ا'نیافاد	'CG'I	'GG	CT'I	GAG	iGAC	ccc	
1		G	G	G	G	D	K	T		_	С		-				Þ	E	L	L	G	•
	121	GG	ACC	GTC	AGT						AAA										ACC	180
	121	CC	TGG	CAG	TCA	AAA	GGA	GAA	GGG	GGG'	TTT	rgg	GTT(CCT	GTGC	GAC	TAC	TAC	AGG	GCC	TGG	
a		G	P	S	V	F	L	F	P	P	K	P	K	D	T	L	M	I	S	R	T	-
	181										CGT								CAAC	TTC	AAC +++	240
	101	GG.	ACT	CCA	GTG	TAC	GCA	CCA	CCA	CCT	GCA	CTC	GGT	GCT'	TCT	GG[ACTO	CAC	TTC	CAAC	TTG	
à		P	E	v	T	С	V	V	v	D	V	S	Н	E	D	P	E	V	K	F	N	•
		TG	GTA	CGI	'GGA	CGG	CGT	GGA	GGT	GCA	TAA'	TGC	CAA	GAC	AAA	3CC(GCG	GA	GGA	GCAC	TAC	200
	241	AC	CAT	GCA	CCI	GCC	GCA	CCT	CCA	CGT.	ATT.	ACG	GTT	CTG	TTT(CGG	CGC	CT	CT	GTO	ATG	300
a		w	Y	v	מ	G	v	E	V	н	N	A	ĸ	T	ĸ	P	R	E	E	Q	Y	-
			CAG	CAC	GTA											CCA	GGA (CTG	CTC	GAAT	rGGC	360
	301	TT	GTC	GTC	CAT						GGA					GGT	CCT	GAC	CGA	CTT	ACCG	300
a		N	s	T	Y	R	V	v	S	V	L	T	v	L	Н	Q	D	W	L	N	G	•
	361										CAA					ccc	CAT	CGA	GAA	AAC	CATC	420
	201	TT														GGG	GTA(GCT	CTT'	rtg(STAG	
a		K	E	Y	K	С	ĸ	V	S	N	K	A	L	P	A	P	I	E	K	T	I	•
	421		CA	AAG	CA						ACC				CAC	CCT	GCC	CCC.	ATC	CCG	GAT	480
	421	AG	GTI	rtc	GTT										GTG	GGA	CGG	GGG	TAG	GGC	CCTA	
a		8	K	A	K	G	Q	P	R	E	P	Q	V	Y	T	L	P	P	Š	R	D	•
	401			rga(CCA	AGAZ	ACC	AGGT	CAG	CCI	GAC	CTG	CCT	GGT	CAA	AGG	CTT	CTA	TCC	CAG	CGAC	540
	481	CI																		GTC	GCTG	,
a		E	L	T	K	N	Q	V	S	L	T	С	L	V	K	G	F	Y	P	S	D	-
	- 4 -	ΑΊ	CGC	CCG	rggi	AGT	3GG/	AGAC	CAA	TGG	GCA	GCC	:GGA	GAA	CAA	CTA	CAA	GAC	CAC	GCC	rccc	600
	541	T	AGC	GGC	ACC:	rca	ccc'	rct	GTI	ACC	CGT	CGC	CCI	CTI	GTT	GAT	GTT	CTG	GTG	CGG	AGGG	
a		I	A	V	E	W	E	s	N	G	Q	Þ	E	N	N	Y	ĸ	T	T	P	P	•

FIG. 26B

	601		• • •		-+-			+				+			-+-			4		• • •	GTCC	660
a		v	Ł	D	S	D	G	s	F	F	L	Y	S	K	L	T	v	D	K	S	R	•
	661				-+-			+				+	, -		-+-			4			CTAC + GATG	720
a		W	Q	Q	G	N	V	F	S	С	S	V	M	Н	E	A	L	Н	N	Н	Y	•
	721		·-		GAG			+	· • • •			+	ATA		GA7		763	3				
•		ጥ	0	ĸ	9	۲.	S	T.	S	P	G	ĸ	*									

SEQUENCE LISTING

<110> LIU, CHUAN-FA
FEIGE, ULRICH
CHEETHAM, JANET
BOONE, THOMAS CHARLES

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ggg gga ccg tca gtc ttc ctc ttc ccc cca aaa ccc aag gac acc ctc 96
Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr Leu
20 25 30

atg atc tcc cgg acc cct gag gtc aca tgc gtg gtg gtg gac gtg agc 144
Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp Val Ser
35 40 45

cac gaa gac cct gag gtc aag ttc aac tgg tac gtg gac ggc gtg gag

His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly Val Glu

50 55 60

gtg cat aat gcc aag aca aag ccg cgg gag gag cag tac aac agc acg 240

65	HIS	Asn	Ala	гÀЗ	70	гÀз	Pro	Arg	GIU	75	GIN	TYL	ASII	ser	80	
					gtc Val											288
					tgc Cys											336
			Thr		tcc Ser											384 -
					cca Pro											432
					gtc Val 150											480
					Gly											528
ccc Pro	gtg Val	ctg Leu	gac Asp 180	tcc Ser	gac Asp	ggc Gly	tcc Ser	ttc Phe 185	ttc Phe	ctc Leu	tac Tyr	agc Ser	aag Lys 190	ctc Leu	acc Thr	576
gtg Val	gac Asp	aag Lys 195	agc Ser	agg Arg	tgg Trp	cag Gln	cag Gln 200	ggg Gly	aac Asn	gtc Val	ttc Phe	tca Ser 205	tgc Cys	tcc Ser	gtg Val	624
atg Met	cat His 210	Glu	gct Ala	ctg Leu	cac	aac Asn 215	cac His	tac Tyr	acg Thr	cag Gln	aag Lys 220	agc Ser	ctc Leu	tcc Ser	ctg Leu	672
			aaa Lys													684
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Gly	Gly	Pro	Ser 20	Val	Phe	Leu	Phe	Pro 25	Pro	Lys	Pro	Lys	Asp 30	Thr	Leu
Met	Ile	Ser 35	Arg	Thr	Pro	G1u	Val 40	Thr	Сув	Val	Val	Val 45	Asp	Val	Ser
His	Glu 50	Asp	Pro	Glu	Val	Lys 55	Phe	Asn	Trp	Tyr	Val 60	Asp	Gly	Val	Glu
Val 65	His	Asn	Ala	Lys	Thr 70	Lys	Pro	Arg	Glu	G1u 75	Gln	Tyr	Asn	Ser	Thr 80
Tyr	Arg	Val	Val	Ser 85	Val	Leu	Thr	Val	Leu 90	His	Gln	Asp	Trp	Leu 95	Asn
Gly	Lys	Glu	Tyr 100	Lys	Cys	Lys	Val	Ser 105	Asn	Lys	Ala	Leu	Pro 110	Ala	Pro
Ile	Glu	Lys 115	Thr	Ile	Ser	Lys	Ala 120	Lys	Gly	Gln	Pro	Arg 125	Glu	Pro	Gln
Val	Tyr 130		Leu	Pro	Pro	Ser 135	Arg	Asp	Glu	Leu	Thr 140	Lys	Asn	Gln	Val
Ser 145	Leu	Thr	Суз	Leu	Val 150	Lys	Gly	Phe	Tyr	Pro 155	Ser	Авр	Ile	Ala	Val 160
Glu	Trp	Glu	Ser	Asn 165	Gly	Gln	Pro	Glu	Asn 170		Tyr	Lys	Thr	Thr 175	Pro
Pro	Val	Leu	Asp 180	Ser	Asp.	. Gly	Ser	Phe 185	Phe	Leu	Tyr	Ser	Lys 190	Leu	Thr
Val	Asp	Lys 195	Ser	Arg	Trp	Gln	Gln 200	Gly	Asn	Val	Phe	Ser 205	Cys	Ser	Val
Met	His 210		Ala	Leu	His	Asn 215		Tyr	Thr	Gln	Lys 220		Leu	Ser	Leu
Ser 225	Pro	Gly	Lys 												_

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                                  10
Arg Ala
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Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala
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cag ccg gag aac aac tac aag acc acg cct ccc gtg ctg gac tcc gac

Gln Pro Glu Asn Asn Tyr Lys Thr Thr Pro Pro Val Leu Asp Ser Asp

160

155

180

ggc tcc ttc ttc ctc tac agc aag ctc acc gtg gac aag agc agg tgg 63.

Gly Ser Phe Phe Leu Tyr Ser Lys Leu Thr Val Asp Lys Ser Arg Trp

185 190 195

175

cag cag ggg aac gtc ttc tca tgc tcc gtg atg cat gag gct ctg cac 680 Gln Gln Gly Asn Val Phe Ser Cys Ser Val Met His Glu Ala Leu His 200 205 210

aac cac tac acg cag aag agc ctc tcc ctg tct ccg ggt aaa ggt gga 728
Asn His Tyr Thr Gln Lys Ser Leu Ser Leu Ser Pro Gly Lys Gly Gly
215 220 225 230

ggt ggt ggt atc gaa ggt ccg act ctg cgt cag tgg ctg gct gct cgt 776
Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg
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gct taatctcgag gatcc 794

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<212> PRT

<213> Artificial Sequence

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Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr Leu 20 25 30

Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp Val Ser 35 40 45

His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly Val Glu 50 55 60

Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn Ser Thr
65 70 75 80

Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln Asp Trp Leu Asn
85 90 95

Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu Pro Ala Pro

100 105 110

Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro Gln
115 120 125

Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn Gln Val 130 135 140

Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala Val-145 150 155 160

Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr Thr Pro 165 170 175

Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu Thr 180 185 190

Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser Val 195 200 205

Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser Leu 210 215 220

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7

-			tgt		-	-	-		_			•		•		104
Суз	Pro	Pro	Cys	Pro	Ala	Pro	Glu		Leu	Gly	Gly	Pro		Val	Phe	
			10					15				•	20			
																160
			cca Pro			_	_			-						152
rea	Pne		PIO	гуя	PIO	гуя	30	THE	теп	Met	TIE	35	ary	THE	PIO	
		25					30					33				
a=a	atc	202	tgc	ata	ata	ata	asc	ata	age	cac	maa	gac.	cct	gag	atc	200
			Cys													
014	40	****	C _J U			45					50					
aaq	ttc	aac	tgg	tac	gtg	gac	ggc	gtg	gag	gtg	cat	aat	gcc	aag	aça	248
			Trp													
55			•	-	60	_	-			65					70	
aag	ccg	cgg	gag	gag	cag	tac	aac	agc	acg	tac	cgt	gtg	gtc	agc	gtc	296
Lys	Pro	Arg	Glu	Glu	Gln	Tyr	Asn	Ser	Thr	Tyr	Arg	Val	Val	Ser	Val	
				75					80					85		
			ctg													344
Leu	Thr	Val	Leu	His	Gln	Asp	Trp	Leu	Asn	Gly	Lys	Glu	Tyr	Lys	Сув	
			90					95					100			
			aac													392
Lys	Val	Ser	Asn	Lys	Ala	Leu	Pro	Ala	Pro	Ile	Glu		Thr	Ile	Ser	
		105					110					115				
		•														440
			ggg													440
Lys		Lys	Gly	Gln	Pro.		Glu	Pro	GIN	vaı		Thr	Leu	PIO	PIO	
	120					125					130					
									 -	200	a+=	300	tac	cta	atc	488
			gag Glu													•••
	Arg	Asp	GIU	reu	140	гåа	ASII	GIII	Val	145	Dea	****	0,0	204	150	
135					140											
	~~~	++0	tat	ccc	200	aac.	atc	acc	ata	σασ	taa	σασ	аσс	aat	aga	536
			Tyr													
פעם	GLY	1116	-3-	155	-	,			160					165	_	
cao	cca	gag	aac	aac	tac	aaσ	acc	acg	cct	ccc	gtg	ctg	gac	tcc	gac	584
Gln	Pro	Glu	Asn	Asn	Tvr	Lys	Thr	Thr	Pro	Pro	Val	Leu	Asp	Ser	Asp	
			170		• -	-		175					180			
ggc	tcc	ttc	ttc	ctc	tac	agc	aag	ctc	acc	gtg	gac	aag	agc	agg	tgg	632
Gly	Ser	Phe	Phe	Leu	Tyr	Ser	Lys	Leu	Thr	Val	Asp	Lys	Ser	Arg	Trp	
-		185			-		190	-				195			_	

_	_	Gly		-									-	Leu		000
		tac Tyr	-	-	_											728
		ggt Gly										_			-	776
		ggt Gly														824
		ctt Leu 265		-		gcat	aato	etc ç	agga	tcc	ī					861
<212 <213	l> 26 2> PF 3> Ar				_		cial	Sequ	ience	e:Fc	· TMP ·	TMP				
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Gly	Gly	Pro		Val	Phe	Leu	Phe	Pro						mb ~	I.eu	
Met			20					25	Pro	Lys	Pro	Lys	Asp 30	IIIL	Dou	
2100	Ile	Ser 35		Thr	Pro	Glu		25					30			
			Arg				Val 40	25 Thr	Cys	Val	Val	Val 45	30 Asp	Val	Ser	
His	Glu 50	35	Arg Pro	Glu	Val	Lys 55	Val 40 Phe	25 Thr Asn	Cys Trp	Val Tyr	Val	Val 45 Asp	30 Asp Gly	Val	Ser Glu	
His Val 65	Glu 50 His	35 Asp	Arg Pro	Glu Lys	Val Thr 70	Lys 55 Lys	Val 40 Phe Pro	25 Thr Asn Arg	Cys Trp Glu	Val Tyr Glu 75	Val Val 60	Val 45 Asp	30 Asp Gly Asn	Val Val Ser	Ser Glu Thr	

Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro Gln 115 120 125

Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn Gln Val 130 135 140

Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala Val 145 150 155 160

Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr Thr Pro 165 170 175

Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu Thr 180 185 190

Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser Val 195 200 205

Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser Leu 210 215 220

Ser Pro Gly Lys Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg 225 230 235 240

Gln Trp Leu Ala Ala Arg Ala Gly Gly Gly Gly Gly Gly Gly Ile 245 250 255

Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg 260 265

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Leu	Arg	Gln		Leu	Ala	Ala	Arg		Gly	Gly	Gly	Gly	Gly	Gly	Gly	
			10					15					20			
ggc	att	gag	ggc	cca	acc	ctt	cgc	caa	tgg	ctt	gca	gca	cgc	gca	ggg	152
						Leu									•	
		25					30					35			-	
~~~	<b>777</b>	aat	<i>aaa</i>	<b>~</b> 2 <b>~</b>	222	act	C2.C	202	+a+	cc=	cct	tac	CC2	<b>~</b> C2	cct	200
				-		Thr			-			-		-		200
-	40	_	_	_	-	45			-		50	-				
						tca Ser										248
55	nea	Leu	GIY	GIŞ	60	Ser	Val	FIIG	пеп	65	FIO	PLO	пуз	PIQ	70	•
•			-			cgg				-		-				296
qaA	Thr	Leu	Met		Ser	Arg	Thr	Pro	Glu 80	Val	Thr	Суз	Val	Val 85	Val	
				75					00					05		
gac	gtg	agc	cac	gaa	gac	cct	gag	gtc	aag	ttc	aac	tgg	tac	gtg	gac	344
Asp	Val	Ser	His	Glu	Asp	Pro	Glu	Val	Lys	Phe	Asn	Trp	Tyr	Val	Asp	
			90					95					100			
aac	ata	asa	ata	cat	aat	gcc	ааσ	aca	aaσ	cca	caa	σаσ	σaσ	cag	tac	392
						Ala										
		105					110					115				
																440
						gtc Val										440
AGII	120	1111	-7-	my	741	125	501				130					
						tac										488
135	ren	ASD	GIY	rys	140	Tyr	гÃа	Cys	гуа	145	Ser	ASII	гуя	MIG	150	
						acc										536
Pro	Ala	Pro	Ile		Lys	Thr	Ile	Ser		Ala	Lys	Gly	Gln		Arg	
				155					160					165		
gaa	cca	cag	ata	tac	acc	ctg	CCC	cca	tcc	cgg	gat	gag	ctg	acc	aag	584
						Leu										
			170					175					180		· men	
							a+-		222	acc	++~	+2+	ccc	מתר.	mac .	- 632
						tgc Cys										J.J.
						•			-	_		_				

190 195 185 680 atc gcc gtg gag tgg gag agc aat ggg cag ccg gag aac aac tac aag Ile Ala Val Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys 200 205 210 acc acg cct ccc gtg ctg gac tcc gac ggc tcc ttc ttc ctc tac agc Thr Thr Pro Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser 215 220 aag ctc acc gtg gac aag agc agg tgg cag cag ggg aac gtc ttc tca 776 Lys Leu Thr Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser 240 235 tgc tcc gtg atg cat gag gct ctg cac aac cac tac acg cag aag agc Cys Ser Val Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser 255 250 855 ctc tcc ctg tct ccg ggt aaa taatggatcc Leu Ser Leu Ser Pro Gly Lys 265 <210> 10 <211> 269 <212> PRT <213> Artificial Sequence <223> Description of Artificial Sequence:TMP-TMP-Fc Met Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly 10 5 Gly Gly Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp . 25 20 Leu Ala Ala Arg Ala Gly Gly Gly Gly Asp Lys Thr His Thr Cys 40 Pro Pro Cys Pro Ala Pro Glu Leu Leu Gly Gly Pro Ser Val Phe Leu 60 55 Phe Pro Pro Lys Pro Lys Asp Thr Leu Met Ile Ser Arg Thr Pro Glu 70 65

85

Val Thr Cys Val Val Val Asp Val Ser His Glu Asp Pro Glu Val Lys

. 90

95

Phe Asn Trp Tyr Val Asp Gly Val Glu Val His Asn Ala Lys Thr Lys
100 105 110

Pro Arg Glu Glu Gln Tyr Asn Ser Thr Tyr Arg Val Val Ser Val Leu 115 120 125

Thr Val Leu His Gln Asp Trp Leu Asn Gly Lys Glu Tyr Lys Cys Lys 130 135 140

Val Ser Asn Lys Ala Leu Pro Ala Pro Ile Glu Lys Thr Ile Ser Lys 145 150 155 160

Ala Lys Gly Gln Pro Arg Glu Pro Gln Val Tyr Thr Leu Pro Pro Ser 165 170 175

Arg Asp Glu Leu Thr Lys Asn Gln Val Ser Leu Thr Cys Leu Val Lys 180 185 190

Gly Phe Tyr Pro Ser Asp Ile Ala Val Glu Trp Glu Ser Asn Gly Gln 195 200 205

Pro Glu Asn Asn Tyr Lys Thr Thr Pro Pro Val Leu Asp Ser Asp Gly 210 215 220

Ser Phe Phe Leu Tyr Ser Lys Leu Thr Val Asp Lys Ser Arg Trp Gln 225 230 235 240

Gln Gly Asn Val Phe Ser Cys Ser Val Met His Glu Ala Leu His Asn 245 250 255

His Tyr Thr Gln Lys Ser Leu Ser Leu Ser Pro Gly Lys 260 265

<210> 11

<211> 789

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:TMP-Fc

<220>

<221> CDS

<222> (39)···. (779)

<400> 11

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_	cgt Arg	_														104
	cac His															152
	gtt Val 40															200
	acc Thr															248
	gag Glu															296
	aag Lys															344
	agc Ser															392
	aag Lys 120						Lys									440
	atc Ile															488
ctg Leu	ccc Pro	cca Pro	tcc Ser	cgg Arg 155	gat Asp	gag Glu	ctg Leu	acc Thr	aag Lys 160	aac Asn	cag Gln	gtc Val	agc Ser	ctg Leu 165	acc Thr	536
tgc Cys	ctg Leu	gtc Val	aaa Lys 170	Gly	ttc Phe	tat Tyr	ccc Pro	agc Şer 175	gac Asp	atc Ile	gcc Ala	gtg Val	gag Glu 180	tgg Trp	gag Glu	584

					gag Glu						-				_	632
					ttc Phe									-	-	680
			_	_	ggg Gly 220		-			-		-	_			728
					tac Tyr							-		_		776
aaa Lys	taat	:ggat	cc			•										789
<211 <212)> 12 l> 24 l> PI	17 RT		- C												
<213 <223					nenc Art		cial	Sequ	ience	: TMI	P-Fc					
<223 <400	3> De 3> 12	escri	iptic	on of	-	ific						Ala	Arg	Ala 15	Gly	
<223 <400 Met 1	3> De 3> 12 Ile	escri ? Glu	iptic Gly	Pro 5	Art	ific	Arg	Gln	Trp 10	Leu	Ala			15		
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<223 <400 Met 1 Gly	3> De 3> 12 Ile Gly Leu	Glu Gly Leu 35	Gly Gly 20 Gly	Pro 5 Asp	Thr Lys	Leu Thr	Arg His Val 40	Gln Thr 25 Phe	Trp 10 Cys Leu	Leu Pro	Ala Pro	Cys Pro 45	Pro 30 Lys	15 Ala Pro	Pro Lys:	
<223 <400 Met 1 Gly Glu Asp	3> De 3> De 3> 12 1le Gly Leu Thr 50	Glu Gly Leu 35	Gly Gly 20 Gly Met	Pro 5 Asp Gly	Thr Lys	Leu Thr Ser	Arg His Val 40	Gln Thr 25 Phe	Trp 10 Cys Leu Glu	Leu Pro Phe Val	Ala Pro Pro Thr 60	Cys Pro 45 Cys	Pro 30 Lys Val	15 Ala Pro Val	Pro Lys	
<223 <400 Met 1 Gly Glu Asp 65	3> De OP 12 Ile Gly Leu Thr 50 Val	Glu Gly Leu 35 Leu Ser	Gly Gly 20 Gly Met	Pro 5 Asp Gly Ile	Thr Lys Pro Ser	Leu Thr Ser Arg 55	Arg His Val 40 Thr	Gln Thr 25 Phe Pro	Trp 10 Cys Leu Glu	Leu Pro Phe Val Phe 75	Ala Pro Pro Thr 60 Asn	Cys Pro 45 Cys	Pro 30 Lys Val	15 Ala Pro Val	Pro Lys Val Asp 80	

Trp Leu Asn Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu 115 120 125

Pro Ala Pro Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg 130 135 140

Glu Pro Gln Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys 145 150 155 160

Asn Gln Val Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp 165 170 175

Ile Ala Val Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys 180 185 190

Thr Thr Pro Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser 195 200 205

Lys Leu Thr Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser 210 225 220

Cys Ser Val Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser 225 230 235 240

Leu Ser Leu Ser Pro Gly Lys
245

<210> 13

<211> 14

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:TMP

<400> 13

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala 1 5 10

<210> 14

<211> 36 ...

<212> PRT

<213> Artificial Sequence

PCT/US99/25044 WO 00/24782

<220> <223> Description of Artificial Sequence: TMP-TMP <400> 14 Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly 10 Gly Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu 25 Ala Ala Arg Ala 35 <210> 15 <211> 812 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:Fc-EMP <220> <221> CDS <222> (39)..(797) <400> 15 totagatttg ttttaactaa ttaaaggagg aataacat atg gac aaa act cac aca 56 Met Asp Lys Thr His Thr tgt cca cct tgt cca gct ccg gaa ctc ctg ggg gga ccg tca gtc ttc 104 Cys Pro Pro Cys Pro Ala Pro Glu Leu Leu Gly Gly Pro Ser Val Phe 15 10 ctc ttc ccc cca aaa ccc aag gac acc ctc atg atc tcc cgg acc cct Leu Phe Pro Pro Lys Pro Lys Asp Thr Leu Met Ile Ser Arg Thr Pro 25 30 gag gtc aca tgc gtg gtg gtg gac gtg ågc cac gaa gac cct gag gtc 200 Glu Val Thr Cys Val Val Val Asp Val Ser His Glu Asp Pro Glu Val 45 40 aag ttc aac tgg tac gtg gac ggc gtg gag gtg cat aat gcc aag aca 248 Lys Phe Asn Trp Tyr Val Asp Gly Val Glu Val His Asn Ala Lys Thr

60

55

65

70

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aag	cca	cac	ı de	ag (gag	cag	ta	са	ac	agc	ac	g t	ac	cgt	gtg	gto	a	дС	gt	2	296
aag Lys	Pro	Arc	G.	lu (Glu	Gln	Ту	r A	sn	Ser	Th:	r T	yr .	Arg	Val	Va]		<u></u>	Va:	l	
-,-					75						8	0						85			
																			. -	_	244
ctc	acc	gto	c c	tg	cac	cag	ga	c t	gg	ctg	aa	t g	gc	aag	gag	tac	: a	ag	tg	С 	344
Leu	Thr	Va:	l L	eu	His	Gln	. As	p 1	(rp	Leu	As	n G	ly.	Lys	Glu	ıy.		ys.	СУ	8	
				90						95						10	J				
																20	. a	+~	to	c	392
aag	gtc	tc	c a	ac	aaa	gco	: ct	C (cca	gcc	CC	C	itc	gag	aaa	mh.	- 1	110	Se	r	
Lys	Val	Se	r A	sn	ГЛа	Ala	ı Le	eu l	Pro	Ala	PI	0 .	rte	GIU	115	111			-	_	
		10	5						110						110						
													~+ <i>~</i>	tac	acc	: ct	ar d	CCC	cc	:a	440
aaa	gco	aa	a g	Igg	cag	CCC	C C	ja '	gaa	CCa	C6	19 Y	y cy Va 1	TVT	Thr	Le	u	Pro	PI	0	
Lys	Ala	Ly	s G	ly	Gln	Pro	O A	rg '	GIU	PIO	G	.11	Val	130				-			
	120)					1.	25													
	cgç									Cac		t C	agc	cto	aco	: tg	rc ·	ctg	gt	c	488
tcc	cgç Arç	g ga	it q	gag 	ctg	ac	c a	ag 	aac aan	Gla	V	al	ser	Lev	Thi	c Cy	78	Leu	V	a l	
		j As	p (31u	Leu	14		ys	VOIT				145						1	50	
135	•					7.4	U														
	r dd					. 20	с п	ac	atc	acc	: g	tg	gag	tgg	ga	gaç	jC	aat	g	gg	536
aaa	gg Gl	ים כ ים	.c '	cat	Dro	. ey	r A	SD.	Ile	Ala	V	al	Glu	TI	G11	u Se	er	Asn	G	1y	
Lys	GI	y Pi	18	īĀī	159	, 5e					1	60						165	•		
	gcc	~ ~:	a /7	aac	aac	c ta	c a	ag	acc	ace	g c	ct	ccc	: gt	g ct	g g	ac	tco	; g	ac	584
ca;	g cc n Pr	o G	111	Asn	Ası	n Ty	r I	ys	Thr	Th:	r P	ro	Pro	va:	l Le	u A	вÞ	Ser	. A	ga.	
GII	I FI	0 0		170				-		17	5					1	80				
																					632
aa	c to	c t	tc	tto	: ct	c ta	ic 8	ıgc	aaq	g ct	c a	CC	gt	g ga	c aa	g a	gc	agg	; t	.gg	032
GJ.	c tc y Se	r P	he	Phe	Le	u Ty	r s	Ber	Lys	s Le	u I	hr	Va!	L As	P -1	-	er	Arq	3 .7	ΤĐ	
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ca	g ca	ıg g	gg	aad	gt	c t	tc	tca	tg	c to	C	gtg	ate	g ca	t ge	19 9 1., 7	12	T.e	9 \ 11 1	lis	
G1	g ca n Gl	n G	ily	Ası	n Va	1 P	he	Ser	Cy	s Se	r 7	Val	·Me	C 113		Lu		20			
	20					•		205	,					21	.0						
												 -		+ ~-	.a a	at ?	aaa	aa	t	gga	728
aa	ic ca	ac t	cac	ac	g ca	ıg a	ag	ago	: ct	c to	:C (ccg		ان با	.U G.	lv I	LVS	Gl	У	Gly	
As	ic ca	is :	lyr	Th	r Gl	ln L	ys	Ser	: Le	u se	SI.	Ten	22			-, .			-	230	
21						2	20						44	_			٠				
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gç	gt g	gt (ggt	gg	a g	gt 8	ct	tac	t C	:	 	uic	, UL	ie G	lv P	ro	Lev	ı Tl	ır	Trp	
G.	jt g ly G	ly (Gly	G1	у G	ly 1	hr	ТУ	ເ 5€	er C	γÞ	240) E1					24	15		
					2	35						~ * ·	-								
										2+~	tco	rt.a	gai	tcc			,				812
g.	tt t	gc	aaa	CC	g c	ag q	gt	gg.	t ti		رب	, ~ 9	2 ~ '							-	
V	al C	ys	Lys			ın (tΖ	GT.	Y												
				25	50																

<210> 16

<211> 253

<212> PRT

<213> Artificial Sequence

<223> Description of Artificial Sequence:Fc-EMP

<400> 16

Met Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu Leu Leu

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Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr Leu 20 25 30

Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp Val Ser 35 40 45

His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly Val Glu 50 55 60

Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Glu Tyr Asn Ser Thr 65 70 75 80

Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln Asp Trp Leu Asn 85 90 95

Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu Pro Ala Pro 100 105 110

Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro Gln 115 120 125

Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn Gln Val 130 135 140

Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala Val 145 150 155 160

Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr Thr Pro 165 170 175

Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu Thr 180 185 190

Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser Val 195 200 205

Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser Leu

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220 215 210

Ser Pro Gly Lys Gly Gly Gly Gly Gly Gly Thr Tyr Ser Cys His 235 230

Phe Gly Pro Leu Thr Trp Val Cys Lys Pro Gln Gly Gly 250 245

<210> 17

<211> 807

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EMP-Fc

<220>

<221> CDS

<222> (39)..(797)

<400> 17

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tgc cac ttc ggc ccg ctg act tgg gta tgt aag cca caa ggg ggt ggg 104 Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys Pro Gln Gly Gly 15 10

gga ggc ggg ggg gac aaa act cac aca tgt cca cct tgc cca gca cct 152 Gly Gly Gly Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro 30 25

gaa ctc ctg ggg gga ccg tca gtt ttc ctc ttc ccc cca aaa ccc aag 200 Glu Leu Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys 45 40

gac acc ctc atg atc tcc cgg acc cct gag gtc aca tgc gtg gtg Asp Thr Leu Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val 65 60 55

gac gtg agc cac gaa gac cct gag gtc aag ttc aac tgg tac gtg gac Asp Val Ser His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp · 80 75

ggc gtg gag gtg cat aat gcc aag aca aag ccg cgg gag gag cag tac 344

Gly	Val	Glu		His	Asn	Ala	Lys		Lys	Pro	Arg	Glu		Gln	Tyr	
			90					95				•	100			
aac	agc	acg	tac	cgt	gtg	gtc	agc	gtc	ctc	acc	gtc	ctg	cac	cag	gac	392
Asn	Ser	Thr	Tyr	Arg	Val	Val	Ser	Val	Leu	Thr	Val	Leu	His	Gln	Asp	
		105					110					115				
taa	cta	aat	aac	aaq	gag	tac	aag	tgc	aag	gtc	tcc	aac	aaa	gcc	ctc	440
Tro	Leu	Asn	Glv	Lvs	Glu	Tyr	Lys	Cys	Lys	Val	Ser	Asn	Lys	Ala	Leu	
	120			-		125	_				130					
cca	acc	CCC	atc	ааа	aaa	acc	atc	tcc	aaa	gcc	aaa	ggg	cag	ccc	cga	488
Pro	Ala	Pro	Ile	Glu	Lys	Thr	Ile	Ser	Lys	Ala	Lys	Gly	Gln	Pro	Arg	
135					140					145					150	
ma a	cca	cag	ata	tac	acc	cta	ccc	cca	tcc	cgg	gat	gag	ctg	acc	aag	536
Glu	Pro	Gln	Val	Tvr	Thr	Leu	Pro	Pro	Ser	Arg	Asp	Glu	Leu	Thr	Lys	
				155					160					165		
aac	cag	gtc	agc	ctg	acc	tgc	ctg	gtc	aaa	ggc	ttc	tat	ccc	agc	gac	584
Asn	Gln	Val	Ser	Leu	Thr	Cys	Leu	Val	Lys	Gly	Phe	Tyr	Pro	Ser	Asp	
			170					175					180			
atc	gcc	gtg	gag	tgg	gag	agc	aat	ggg	cag	ccg	gag	aac	aac	tac	aag	632
Ile	Ala	Val	Glu	Trp	Glu	Ser	Asn	Gly	Gln	Pro	Glu	Asn	Asn	Tyr	Lys	
		185					190	-				195				•
acc	acg	cct	ccc	gtg:	ctg	gac	tcc	gac	ggc	tcc	ttc	ttc	cto	tac	agc	680
Thr	Thr	Pro	Pro	Val	Leu	Asp	Ser	Asp	Gly	Ser	Phe	Phe	Lev	Tyr	Ser	
	200					205					210					
aag	cto	acc	: gtg	gac	: aaç	ago	agg:	tgg	cag	cag	ggg	aac	gto	tto	tca	728
Lys	Leu	Thr	. Val	Asr	Lys	Ser	Arg	Tr	Gln	Gln	Gly	Asn	Va]	Phe	ser	
215					220					225			:		230	
tgo	tco	gte	, atq	, cat	gaç	g gct	ctg	cac	aac	cac	tac	acç	caq	y aag	agc	776
Сув	Sei	. Val	L Met	. His	Gli	ı Ala	a Lev	His	Asn	1 His	туг	Thi	Gli	n Lys	s ser	
		•		235					240)				245	•	
cto	: tc	c cto	g to	t cc	g ggt	t aaa	a taa	tgga	atcc		as a					807
		r Le														
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<210> 18
<211> 253
<212> PRT
<213> Artificial Sequence

PCT/US99/25044 WO 00/24782

<223> Description of Artificial Sequence: EMP-Fc

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- Met Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys
- Lys Pro Gln Gly Gly Gly Gly Gly Gly Asp Lys Thr His Thr Cys
- Pro Pro Cys Pro Ala Pro Glu Leu Leu Gly Gly Pro Ser Val Phe Leu
- Phe Pro Pro Lys Pro Lys Asp Thr Leu Met Ile Ser Arg Thr Pro Glu
- Val Thr Cys Val Val Val Asp Val Ser His Glu Asp Pro Glu Val Lys
- Phe Asn Trp Tyr Val Asp Gly Val Glu Val His Asn Ala Lys Thr Lys
- Pro Arg Glu Glu Gln Tyr Asn Ser Thr Tyr Arg Val Val Ser Val Leu
- Thr Val Leu His Gln Asp Trp Leu Asn Gly Lys Glu Tyr Lys Cys Lys
- Val Ser Asn Lys Ala Leu Pro Ala Pro Ile Glu Lys Thr Ile Ser Lys
- Ala Lys Gly Gln Pro Arg Glu Pro Gln Val Tyr Thr Leu Pro Pro Ser
- Arg Asp Glu Leu Thr Lys Asn Gln Val Ser Leu Thr Cys Leu Val Lys
- Gly Phe Tyr Pro Ser Asp Ile Ala Val Glu Trp Glu Ser Asn Gly Gln
- Pro Glu Asn Asn Tyr Lys Thr Thr Pro Pro Val Leu Asp Ser Asp Gly
- Ser Phe Phe Leu Tyr Ser Lys Leu Thr Val Asp Lys Ser Arg Trp Gln
- Gln Gly Asn Val Phe Ser Cys Ser Val Met His Glu Ala Leu His Asn

His Tyr Thr Gln Lys Ser Leu Ser Leu Ser Pro Gly Lys 245

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210>	19															
211>																
:212>																
			cial	Semi	ance	,									-	
213	ALU	.1110	.TaI	sequ	ence	•										
<220>	•				* 4- 4	£1.4	-1 0	20011	מחתם	EMP.	EMP-	Fc				
<223>	> Des	cri	ption	OI	ALCI	LICI	.aı .	equ								
<220	>															
_	> CDS															
<222	> (4:	1)	(871)													
•																
<400	> 19												-~-	• ~ +	+20	55
tcta	gatt	tg a	gtttl	taact	t tt	tagaa	agga	gga	ataa	aat	atg (gya (71	mh-	mr.	
											Met (GIY (этХ	THE		
											1				5	
tct	tac	cac	ttc	ggc	cca	ctg	act	tgg	gtt	tgc	aaa	ccg	cag	ggt	ggc	103
Ser	Cvs	His	Phe	Glv	Pro	Leu	Thr	Trp	Val	Суз	Lys	Pro	Gln	Gly	Gly	
D C2	-,-			10					15					20		
		~~~	ggc	aat	aat	acc	tat	tcc	tgt	cat	ttt	ggc	ccg	ctg	acc	151
ggc	ggc	gge	Gly	990 010	99°	Thr	ጥሆን	Ser	Cvs	His	Phe	Gly	Pro	Leu	Thr	
GIĀ	GIY	GIA		GIŞ	GIJ	****	-3-	30			,		35			
			25					•								
			aag				~~+	aaa	aaa	aac	aaa	aaa	gac	aaa	act	199
tgg	gta	tgt	aag Lys	CCA	Caa	999	99 c	333	G1v	Glv	Glv	Glv	Asp	Lys	Thr	
Trp	Val		Lys	Pro	GIN	GIY	GIA	GIY	GIY	017		50		-		
		40					45					-				
										a+ a	ata	aaa	aga	cca	tca	247
cac	aca	tgt	сса	cct	tgc	cca	gca	CCE	gaa	* ***	Tou	G1 17	Glv	Pro	Ser	
His	Thr	Cys	Pro	Pro	Cys	Pro	Ala	Pro	GIU	Leu	Tierr	GIY	GIJ			
	55					60					65					
														+00	caa	295
gtt	ttc	ctc	ttc	CCC	cca	aaa	CCC	aag	gac	acc	ctc	atg	atc	7	cgg	2,5
Val	Phe	Leu	Phe	Pro	Pro	Lys	Pro	ГЛа	Asp	Thr	Leu	Met	116	s Ser	**** 4	
70					75					80					85	
																242
acc	cct	σaσ	atc	aca	tgc	gtg	gtg	gtg	gac	gtg	agc	cac	gaa	gac	cct Pro	343
ጥኮታ	Pro	G1u	Val	Thr	Cys	Val	Val	Val	Asp	Val	Ser	His	Glu			
				90					95	,			***	100	)	
				-			•								-	
		. ,,,,	. ++~	aac	taa	tac	gta	gad	ggc	gto	gag	gtg	cat	t aat	gcc Ala	391
gag	, 47.3	, aaç	, ccc	\ \an	مديل ۾ ھي	TV	Val	Ast	Gly	Val	Glu	Val	Hi	s Ası	a Ala	
تالمن	val	. ⊔y≿	, 5116	47011	. ~	-3-		-	_							

115 110 105 aag aca aag ccg cgg gag gag cag tac aac agc acg tac cgt gtg gtc Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn Ser Thr Tyr Arg Val Val 125 120 age gte etc ace gte etg cae cag gae tgg etg aat gge aag gag tae Ser Val Leu Thr Val Leu His Gln Asp Trp Leu Asn Gly Lys Glu Tyr 140 aag tgc aag gtc tcc aac aaa gcc ctc cca gcc ccc atc gag aaa acc Lys Cys Lys Val Ser Asn Lys Ala Leu Pro Ala Pro Ile Glu Lys Thr 160 155 150 atc tcc aaa gcc aaa ggg cag ccc cga gaa cca cag gtg tac acc ctg 583 Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro Gln Val Tyr Thr Leu 175 170 ecc cca tcc egg gat gag etg acc aag aac cag gte age etg ace tge Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn Gln Val Ser Leu Thr Cys 190 185 ctg gtc aaa ggc ttc tat ccc agc gac atc gcc gtg gag tgg gag agc 679 Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala Val Glu Trp Glu Ser 210 205 200 aat ggg cag ccg gag aac aac tac aag acc acg cct ccc gtg ctg gac 727 Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr Thr Pro Pro Val Leu Asp 225 220 215 tcc gac ggc tcc ttc ttc ctc tac agc aag ctc acc gtg gac aag agc 775 Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu Thr Val Asp Lys Ser 240 235 230 agg tgg cag cag ggg aac gtc ttc tca tgc tcc gtg atg cat gag gct Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser Val Met His Glu Ala 255 250 ctg cac aac cac tac acg cag aag agc ctc tcc ctg tct ccg ggt aaa Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser Leu Ser Pro Gly Lys 275 270 265

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<210> 20 <211> 277 <212> PRT

<213> Artificial Sequence

<223> Description of Artificial Sequence: EMP-EMP-Fc

<400> 20

- Met Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys

  1 5 10 15
- Lys Pro Gln Gly Gly Gly Gly Gly Gly Gly Thr Tyr Ser Cys His
  20 25 30
- Phe Gly Pro Leu Thr Trp Val Cys Lys Pro Gln Gly Gly Gly Gly Gly 35 40 45
- Gly Gly Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu Leu 50 55 60
- Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr 65 70 75 80
- Leu Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp Val 85 90 95
- Ser His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly Val 100 105 110
- Glu Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn Ser 115 120 125
- Thr Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln Asp Trp Leu 130 135 140
- Asn Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu Pro Ala 145 150 155 160
- Pro Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro 165 170 175
- Gln Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn Gln
  180 185 190
  - Val Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala 195 200 205
  - Val Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr Thr 210 215 220
  - Pro Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu 225 230 235 240

PCT/US99/25044 WO 00/24782

Thr Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser

250 245 Val Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser 270 265 Leu Ser Pro Gly Lys 275 <210> 21 <211> 884 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:Fc-EMP-EMP <220> <221> CDS <222> (39)..(869) <400> 21 tctagatttg ttttaactaa ttaaaggagg aataacat atg gac aaa act cac aca 56 Met Asp Lys Thr His Thr 1 tgt cca cct tgc cca gca cct gaa ctc ctg ggg gga ccg tca gtt ttc Cys Pro Pro Cys Pro Ala Pro Glu Leu Leu Gly Gly Pro Ser Val Phe 15 10 ctc ttc ccc cca aaa ccc aag gac acc ctc atg atc tcc cgg acc cct 152 Leu Phe Pro Pro Lys Pro Lys Asp Thr Leu Met Ile Ser Arg Thr Pro 25 30 gag gtc aca tgc gtg gtg gtg gac gtg agc cac gaa gac cct gag gtc 200 Glu Val Thr Cys Val Val Val Asp Val Ser His Glu Asp Pro Glu Val 45 40 aag ttc aac tgg tac gtg gac ggc gtg gag gtg cat aat gcc aag aca 248 Lys Phe Asn Trp Tyr Val Asp Gly Val Glu Val His Asn Ala Lys Thr 65 60 55

75

aag ccg cgg gag gag cag tac aac agc acg tac cgt gtg gtc agc gtc Lys Pro Arg Glu Glu Gln Tyr Asn Ser Thr Tyr Arg Val Val Ser Val

80

85

		gtc														344
Leu	Thr	Val	Leu 90	His	Gln	Asp	Trp	Leu 95	Asn	Gly	Lys	Glu	Tyr 100	Lys	Cys	
aag	gtc	tcc	aac	aaa	gcc	ctc	cca	gcc	ccc	atc	gag	aaa	acc	atc	tcc	392
Lys	Val	Ser	Asn	Lys	Ala	Leu	Pro	Ala	Pro	Ile	Glu	Lys	Thr	Ile	Ser	
		105					110					115				
		aaa														440
Lys	Ala	Lys	Gly	Gln	Pro	Arg	Glu	Pro	Gln	Val		Thr	Leu	Pro	Pro	
	120					125					130					
		gat'														488
Ser	Arg	Asp	Glu	Leu		rys	Asn	Gln	Val		Leu	Thr	Cys	Leu		
135					140					145					150	
		ttc														536
Lys	Gly	Phe	Tyr		Ser	qaA	Ile	Ala		Glu	Trp	Glu	Ser		Gly	
				155					160					165		
		gag														584
Gln	Pro	Glu	Asn	Asn	Tyr	Lys	Thr	Thr	Pro	Pro	Val			Ser	Asp	
			170					175					180			
		ttc														632
Gly	Ser	Phe	Phe	Leu	Tyr	Ser	Lys	Leu	Thr	Val	Asp		Ser	Arg	Trp	
		185					190					195				
		ggg														680
Gln	Gln	Gly	Asn	Val	Phe	Ser	Cys	Ser	Val	Met		Glu	Ala	Leu	His	
	200					205					210					
aac	cac	tac	acg	cag	aag	agc	ctc	tcc	ctg	tct	ccg	ggt	aaa	ggt	gga	728
Asn	His	Tyr	Thr	Gln	Lys	. Ser	Leu	Ser	Leu		Pro	Gly	Lys	Gly		
215					220					225				•	230	
ggt	ggt	ggc	gga	ggt	act	tac	tct	tgc	cac	ttc	ggc	cca	ctg	act	tgg	776
Gly	Gly	Gly	Gly	Gly	Thr	Tyr	Ser	Сла		Phe	Gly	Pro	Leu	Thr	Trp	
				235					240					245		
gtt	tgc	aaa	ccg	cag	ggt	ggc	ggc	ggc	ggc	ggc	ggt	ggt	acc	tat	tcc	824
Val	Суз	Lys	Pro	Gln	Gly	Gly	Gly			Gly	Gly	Gly	Thr	Туг	Ser	
			250					255					260			
tgt	cat	ttt	 ggc	ccg	ctg	acc	tgg	ġta	tgt	aag	сса	caa	ggg	ggt	<b>.</b> -	869
Сув	His	Phe		Pro	Leu	Thr	Trp	Val	Суз	ГЛа	Pro	Gln	Gly	Gly	•	
		265					270					275	)			

taatctcgag gatcc

884

- <210> 22
- <211> 277
- <212> PRT
- <213> Artificial Sequence
- <223> Description of Artificial Sequence:Fc-EMP-EMP

<400> 22

- Met Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu Leu Leu 1 5 10 15
- Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr Leu 20 25 30
- Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp Val Ser 35 40 45
- His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly Val Glu 50 55 60
- Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn Ser Thr 65 70 75 80
- Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln Asp Trp Leu Asn 85 90 95
- Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu Pro Ala Pro 100 105 110
- Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro Gln 115 120 125
- Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn Gln Val 130 135 140
- Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala Val 145 150 155 160
- Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr Thr Pro 165 170 175
- Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu Thr
- Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser Val

195 200 205

Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser Leu 210 215 220

Ser Pro Gly Lys Gly Gly Gly Gly Gly Gly Thr Tyr Ser Cys His 225 230 235 240

Phe Gly Pro Leu Thr Trp Val Cys Lys Pro Gln Gly Gly Gly Gly Gly 255

Gly Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys 260 265 270

Lys Pro Gln Gly Gly 275

<210> 23

<211> 1545

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:pAMG216

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ctcgaggatc cgcggaaaga agaagaagaa gaagaaagcc cgaaaggaag ctgagttggc 1440
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<210> 24
<211> 14
<212> PRT
<213> Artificial Sequence
.<220>
<223> Description of Artificial Sequence: TPO-MIMETIC
       PEPTIDE
<400> 24
Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Lys Ala
                  5
                                     10
  1
 <210> 25
 <211> 14
<212> PRT
<213> Artificial Sequence
<220>
 <223> Description of Artificial Sequence: TPO-MIMETIC
       PEPTIDE
 <400> 25
Ile Glu Gly Pro Thr Leu Arg Glu Trp Leu Ala Ala Arg Ala
                                    10
                 5
<210> 26
 <211> 29
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: TPO-MIMETIC
      PEPTIDE
 <220>
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<223> At position 15, Xaa=a linker sequence of 1 to 20 amino acids

<400> 26

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Xaa Ile 1. 5 10 15

Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala
20 25

<210> 27

<211> 29

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

<220>

<223> At position 15, Xaa=a linker sequence of 1 to 20 amino acids

<400> 27

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Lys Ala Xaa Ile
1 5 10 15

Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Lys Ala 20 25

<210> 28

<211> 29

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO·MIMETIC PEPTIDE

<220>

<223> At position 9 disulfide linkage with residue 24

<220>

<223> At position 24 disulfide linkage with residue 9

<400> 28 Ile Glu Gly Pro Thr Leu Arg Gln Cys Leu Ala Ala Arg Ala Xaa Ile 1 5 Glu Gly Pro Thr Leu Arg Gln Cys Leu Ala Ala Arg Ala 20 <210> 29 <211> 31 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE <220> <223> At position 16 bromoacetyl group linked to sidechain <400> 29 Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Xaa Lys 5 Xaa Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala 25 20 <210> 30 <211> 31 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE <223> At position 16 polyethylene glycol linked to sidechain

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Xaa Lys

10

Xaa Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala 20 25 30

<210> 31

<211> 29

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:TPO-MIMETIC PEPTIDE

<220>

<223> At position 9 disulfide bond to residue 9 of a separate identical sequence

<400> 31

Ile Glu Gly Pro Thr Leu Arg Gln Cys Leu Ala Ala Arg Ala Xaa Ile 1 5 10 15

Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala 20 25

<210> 32

<211> 29

<212> PRT

<213> Artificial Sequence

<220>

<220>

<223> At position 24 disulfide bond to residue 9 of a separate identical sequence

<400> 32

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Xaa Ile
1 5 10 15

Glu Gly Prö Thr Leu Arg Gln Cys Leu Ala Ala Arg Ala
20 25

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<210> 33
<211> 9
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: TPO-MIMETIC
      PEPTIDE
<400> 33
Val Arg Asp Gln Ile Xaa Xaa Xaa Leu
<210> 34
<211> 6
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TPO-MIMETIC
      PEPTIDE
<400> 34
Thr Leu Arg Glu Trp Leu
 1 . 5
 <210> 35
 <211> 10
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: TPO-MIMETIC
      PEPTIDE
 <400> 35
 Gly Arg Val Arg Asp Gln Val Ala Gly Trp
                 5 10
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<210> 36

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<211> 10
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: TPO-MIMETIC
      PEPTIDE
<400> 36
Gly Arg Val Lys Asp Gln Ile Ala Gln Leu
                  5
<210> 37
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:Description of
      Artificial SequenceTPO-MIMETIC PEPTIDE
<400> 37
Gly Val Arg Asp Gln Val Ser Trp Ala Leu
<210> 38
<211> 10
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: TPO-MIMETIC
      PEPTIDE
<400> 38
Glu Ser Val Arg Glu Gln Val Met Lys Tyr
                  5
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<210> 39 <211> 10 <212> PRT <213> Artificial Sequence

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<220>
 <223> Description of Artificial Sequence: TPO-MIMETIC
 <400> 39
 Ser Val Arg Ser Gln Ile Ser Ala Ser Leu
                 5
 <210> 40
 <211> 10
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: TPO-MIMETIC
      PEPTIDE
 <400> 40
 Gly Val Arg Glu Thr Val Tyr Arg His Met
  1 5
<210> 41
 <211> 11
 <212> PRT
<213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: INTEGRIN
      BINDING PEPTIDE
 <400> 41
 Gly Val Arg Glu Val Ile Val Met His Met Leu
  1
         5
 <210> 42
 <211> 11
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: TPO-MIMETIC
```

PEPTIDE

<400> 42
Gly Arg Val Arg Asp Gln Ile Trp Ala Ala Leu
1 5 10

<210> 43

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

<400> 43

Ala Gly Val Arg Asp Gln Ile Leu Ile Trp Leu

1 5 10

<210> 44

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

<400> 44

Gly Arg Val Arg Asp Gln Ile Met Leu Ser Leu

1 5 10

<210> 45

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

<400> 45

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Gly Arg Val Arg Asp Gln Ile Xaa Xaa Xaa Leu
1 5 10
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<210> 46

<211> 10

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:TPO-MIMETIC
 PEPTIDE

<400> 46

Cys Thr Leu Arg Gln Trp Leu Gln Gly Cys
1 5 10

<210> 47

<211> 10

<212> PRT

<213> Artificial Sequence

<220>

<400> 47

Cys Thr Leu Gln Glu Phe Leu Glu Gly Cys
1 5 10

<210> 48

<211> 10

<212> PRT

<213> Artificial Sequence

<220>

<400> 48

Cys Thr Arg Thr Glu Trp Leu His Gly Cys
1 5 10

```
<210> 49
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TPO-MIMETIC
      PEPTIDE
<400> 49
Cys Thr Leu Arg Glu Trp Leu His Gly Gly Phe Cys
  1
                  5
<210> 50
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:Fc-TMP
<400> 50
Cys Thr Leu Arg Glu Trp Val Phe Ala Gly Leu Cys
                                     10
<210> 51
<211> 13
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:Fc-TMP
 <400> 51
 Cys Thr Leu Arg Gln Trp Leu Ile Leu Leu Gly Met Cys
                                      10
                   5
  1
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<210> 52 ... <211> 14 <212> PRT <u>`</u>^.

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<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TPO-MIMETIC
      PEPTIDE
<400> 52
Cys Thr Leu Ala Glu Phe Leu Ala Ser Gly Val Glu Gln Cys
                                    10
  1
                 5
<210> 53
<211> 14
<212> PRT
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<220>
<223> Description of Artificial Sequence:Fc-TMP
<400> 53
Cys Ser Leu Gln Glu Phe Leu Ser His Gly Gly Tyr Val Cys
                                    10
<210> 54
<211> 14
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:Fc-TMP
<400> 54
Cys Thr Leu Arg Glu Phe Leu Asp Pro Thr Thr Ala Val Cys
                                     10
                  5
<210> 55
<211> 14
<212> PRT
<213> Artificial Sequence
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<220>
<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

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<400> 55
Cys Thr Leu Lys Glu Trp Leu Val Ser His Glu Val Trp Cys
1 5 10
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<210> 56

<211> 10

<212> PRT

<213> Artificial Sequence

<220>

<400> 56

Cys Thr Leu Arg Glu Trp Leu Xaa Xaa Cys
1 5 10

<210> 57

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

<400> 57

Cys Thr Leu Arg Glu Trp Leu Xaa Xaa Xaa Cys 1 5 10

<210> 58

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<400> 58

Cys Thr Leu Arg Glu Trp Leu Xaa Xaa Xaa Cys

1 5 10

<210> 59

<211> 13

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:TPO-MIMETIC
 PEPTIDE

<400> 59

Cys Thr Leu Arg Glu Trp Leu Xaa Xaa Xaa Xaa Xaa Cys

<210> 60

<211> 14

<212> PRT

<213> Artificial Sequence

/22A>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

<400> 60

Cys Thr Leu Arg Glu Trp Leu Xaa Xaa Xaa Xaa Xaa Xaa Cys

<210> 61

<211> 10

<212> PRT

<213> Artificial Sequence

<220>

<400> 61

Arg Glu Gly Pro Thr Leu Arg Gln Trp Met

1 5 10

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<210> 62
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:TPO-MIMETIC
     PEPTIDE
Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala
1 5
<210> 63
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TPO-MIMETIC
     PEPTIDE
<400> 63
Glu Arg Gly Pro Phe Trp Ala Lys Ala Cys
1 5
<210> 64
<211> 10
<212> PRT
<213> Artificial Sequence
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     PEPTIDE
Arg Glu Gly Pro Arg Cys Val Met Trp Met
     5
1
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<210> 65 <211> 14

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<212> PRT
<213> Art
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<213> Artificial Sequence

<220>

<400> 65

Cys Gly Thr Glu Gly Pro Thr Leu Ser Thr Trp Leu Asp Cys
1 5 10

<210> 66

<211> 14

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

<400> 66

Cys Glu Gln Asp Gly Pro Thr Leu Leu Glu Trp Leu Lys Cys
1 5 10

<210> 67

<211> 14

<212> PRT

<213> Artificial Sequence

<220>

<400> 67

Cys Glu Leu Val Gly Pro Ser Leu Met Ser Trp Leu Thr Cys

1 5 10

<210> 68

<211> 14

<212> PRT ...

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

<400> 68

Cys Leu Thr Gly Pro Phe Val Thr Gln Trp Leu Tyr Glu Cys
1 5 10

<210> 69

<211> 14

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

<400> 69

Cys Arg Ala Gly Pro Thr Leu Leu Glu Trp Leu Thr Leu Cys
1 5 10

<210> 70

<211> 14

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

<400> 70

Cys Ala Asp Gly Pro Thr Leu Arg Glu Trp Ile Ser Phe Cys
1 5 10

<210> 71

<211> 13

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

```
<400> 71
Cys Xaa Glu Gly Pro Thr Leu Arg Glu Trp Leu Xaa Cys
                5
<210> 72
<211> 14
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TPO-MIMETIC
      PEPTIDE
<400> 72
Cys Xaa Xaa Glu Gly Pro Thr Leu Arg Glu Trp Leu Xaa Cys
                                    10
                 5
<210> 73
<211> 14
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TPO-MIMETIC
      PEPTIDE
<400> 73
Cys Xaa Glu Gly Pro Thr Leu Arg Glu Trp Leu Xaa Xaa Cys
                 5
                                    10
<210> 74
<211> 15
<212> PRT
<213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: TPO-MIMETIC
      PEPTIDE
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Cys Xaa Xaa Glu Gly Pro Thr Leu Arg Glu Trp Leu Xaa Xaa Cys

1 5 10 15

<210> 75

<211> 16

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

<400> 75

Gly Gly Cys Thr Leu Arg Glu Trp Leu His Gly Gly Phe Cys Gly Gly
1 5 10 15

<210> 76

<211> 18

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:TPO-MIMETIC
 PEPTIDE

<400> 76

Gly Gly Cys Ala Asp Gly Pro Thr Leu Arg Glu Trp Ile Ser Phe Cys
1 5 10 15

Gly Gly

<210> 77

<211> 19

<212> PRT

<213> Artificial Sequence

<220>

<400> 77

Gly Asn Ala Asp Gly Pro Thr Leu Arg Gln Trp Leu Glu Gly Arg Arg

1 5 10 15

Pro Lys Asn

<210> 78

<211> 19

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:TPO-MIMETIC PEPTIDE

<400> 78

Leu Ala Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu His Gly Asn Gly

1 5 10 15

Arg Asp Thr

<210> 79

<211> 19

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

<400> 79

His Gly Arg Val Gly Pro Thr Leu Arg Glu Trp Lys Thr Gln Val Ala

Thr Lys Lys

<210> 80

<211> 18

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:TPO-MIMETIC
 PEPTIDE

<400> 80

Thr Ile Lys Gly Pro Thr Leu Arg Gln Trp Leu Lys Ser Arg Glu His 1 5 10 15

Thr Ser

<210> 81

<211> 18

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:TPO-MIMETIC
 PEPTIDE

<400> 81

Ile Ser Asp Gly Pro Thr Leu Lys Glu Trp Leu Ser Val Thr Arg Gly
1 5 10 15

Ala Ser

<210> 82

<211> 18

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:TPO-MIMETIC
 PEPTIDE

<400> 82

Ser Ile Glu Gly Pro Thr Leu Arg Glu Trp Leu Thr Ser Arg Thr Pro 1 5 10 15

His Ser

```
<210> 83
<211> 14
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EPO-MIMETIC
      PEPTIDE
<400> 83
Tyr Xaa Cys Xaa Xaa Gly Pro Xaa Thr Trp Xaa Cys Xaa Pro
           5
<210> 84
<211> 28
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EPO-MIMETIC
      PEPTIDE
<400> 84
Tyr Xaa Cys Xaa Xaa Gly Pro Xaa Thr Trp Xaa Cys Xaa Pro Tyr Xaa
                                                        15
                5
Cys Xaa Xaa Gly Pro Xaa Thr Trp Xaa Cys Xaa Pro
                                25
             20
<210> 85
<211> 29
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: EPO-MIMETIC
      PEPTIDE
 <220>
 <223> At position 15, Xaa=a linker sequence of 1 to 20
       amino acids
```

<400> 85

```
Tyr Xaa Cys Xaa Xaa Gly Pro Xaa Thr Trp Xaa Cys Xaa Pro Xaa Tyr
1 5 10 15
```

Xaa Cys Xaa Xaa Gly Pro Xaa Thr Trp Xaa Cys Xaa Pro 20 25

<210> 86

<211> 14

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:EPO-MIMETIC
 PEPTIDE

<220>

<223> At position 15 linked through epsilon amine to lysyl, which is linked to a separate identical sequence through that sequence's alpha amine

<400> 86

Tyr Xaa Cys Xaa Xaa Gly Pro Xaa Thr Trp Xaa Cys Xaa Pro 1 5 10

<210> 87

<211> 20

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EPO-MIMETIC PEPTIDE

<400> 87

Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys

1 10 15

Pro Gln Gly Gly

20

<210> 88

<211> 20

```
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EPO-MIMETIC
     PEPTIDE
<400> 88
Gly Gly Asp Tyr His Cys Arg Met Gly Pro Leu Thr Trp Val Cys Lys
                        10
               5
Pro Leu Gly Gly
            20
<210> 89
<211> 20
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: EPO-MIMETIC
      PEPTIDE
<400> 89
Gly Gly Val Tyr Ala Cys Arg Met Gly Pro Ile Thr Trp Val Cys Ser
                                                    15
         5
Pro Leu Gly Gly
             20
 <210> 90
 <211> 20
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: EPO-MIMETIC
      PEPTIDE
 <400> 90
 Val Gly Asn Tyr Met Cys His Phe Gly Pro Ile Thr Trp Val Cys Arg
  1 ... 5 . 10
```

Pro Gly Gly Gly

20

<210> 91 <211> 20

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:EPO-MIMETIC
 PEPTIDE

<400> 91

Gly Gly Leu Tyr Leu Cys Arg Phe Gly Pro Val Thr Trp Asp Cys Gly
1 5 10 15

Tyr Lys Gly Gly 20

<210> 92

<211> 40

<212> PRT

<213> Artificial Sequence

<220>

<400> 92

Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys
1 5 10 15

Pro Gln Gly Gly Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr 20 25 30

Trp Val Cys Lys Pro Gln Gly Gly
35 40

<210> 93

<211> 41

<212> PRT ...

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EPO-MIMETIC PEPTIDE

<220>

<223> At position 21, Xaa=a linker sequence of 1 to 20 amino acids

Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys 10 5

Pro Gln Gly Gly Kaa Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu 25 20

Thr Trp Val Cys Lys Pro Gln Gly Gly 40 35

<210> 94

<211> 23

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EPO-MIMETIC PEPTIDE

<400> 94

Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys 10 5

Pro Gln Gly Gly Ser Ser Lys 20

<210> 95

<211> 46

<212> PRT

<213> Artificial Sequence

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDE

<400> 95

Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys
1 5 10 15

Pro Gln Gly Gly Ser Ser Lys Gly Gly Thr Tyr Ser Cys His Phe Gly 20 25 30

Pro Leu Thr Trp Val Cys Lys Pro Gln Gly Gly Ser Ser Lys 35 40 45

<210> 96

<211> 47

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:TPO-MIMETIC PEPTIDE

<220>

<223> At position 24, Xaa=a linker sequence of 1 to 20 amino acids

<400> 96

Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys
1 5 10 15

Pro Gln Gly Gly Ser Ser Lys Xaa Gly Gly Thr Tyr Ser Cys His Phe 20 25 30

Gly Pro Leu Thr Trp Val Cys Lys Pro Gln Gly Gly Ser Ser Lys
35 40 45

<210> 97

<211> 22

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EPO-MIMETIC PEPTIDE

<220>

<223> At position 22 linked through epsilon amine to lysyl, which is linked to a separate identical

sequence through that sequence's alpha amine

<400> 97

Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys
1 5 10 15

Pro Gln Gly Gly Ser Ser 20

<210> 98

<211> 23

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EPO-MIMETIC PEPTIDE

<220>

<223> At position 23 biotin linked to the sidechain through a linker

<400> 98

Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys
1 5 10 15

Pro Gln Gly Gly Ser Ser Lys
20

<210> 99

<211> 5

<212> PRT

<213> Artificial Sequence

<220>

<220>

<223> At position 4 disulfide bond to residue 4 of a separate identical sequence

<400> 99

Glu Glu Asp Cys Lys

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1 5

<210> 100

<211> 5

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:G-CSF MIMETIC PEPTIDE

<220>

<223> At position 4, Xaa is an isoteric ethylene spacer linked to a separate identical sequence

<400> 100

Glu Glu Asp Xaa Lys 1

<210> 101

<211> 5

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:G-CSF MIMETIC PEPTIDE

<220>

<223> At position 1, Xaa is a pyroglutamic acid residue

<220>

<223> At position 4, Xaa is an isoteric ethylene spacer linked to a separate identical sequence

<400> 101

Xaa Glu Asp Xaa Lys

<210> 102 ....

<211> 5

<212> PRT

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<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EPO-MIMETIC
      PEPTIDE
<220>
<223> At position 1, Xaa is a picolinic acid residue
<220>
<223> At position 4, Xaa is an isoteric ethylene spacer
      linked to a separate identical sequence
<400> 102
Xaa Ser Asp Xaa Lys
  1
<210> 103
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EPO-MIMETIC
      PEPTIDE
<220>
<223> At position 6, Xaa=a linker sequence of 1 to 20
      amino acids
<400> 103
Glu Glu Asp Cys Lys Xaa Glu Glu Asp Cys Lys
                 5
<210> 104
<211> 11
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: EPO-MIMETIC
```

<220>

PEPTIDE

```
<223> At position 6, Xaa=a linker sequence of 1 to 20
      amino acids
<400> 104
Glu Glu Asp Xaa Lys Xaa Glu Glu Asp Xaa Lys
     5
<210> 105
<211> 6
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: ANTIVIRAL (HBV)
     PEPTIDE
<400> 105
Leu Leu Gly Arg Met Lys
1
                5
<210> 106
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TNF-ANTAGONIST
      PEPTIDE
<400> 106
Tyr Cys Phe Thr Ala Ser Glu Asn His Cys Tyr
                  5
                                   10
 1
<210> 107
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TNF-ANTAGONIST
      PEPTIDE
```

```
<400> 107
Tyr Cys Phe Thr Asn Ser Glu Asn His Cys Tyr
                 5
<210> 108
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TNF-ANTAGONIST
      PEPTIDE
<400> 108
Tyr Cys Phe Thr Arg Ser Glu Asn His Cys Tyr
        5
<210> 109
<211> 9
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TNF-ANTAGONIST
      PEPTIDE
<400> 109
Phe Cys Ala Ser Glu Asn His Cys Tyr
<210> 110
<211> 9
<212> PRT
<213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: TNF-ANTAGONSIT
      PEPTIDE
 <400> 110 ...
 Tyr Cys Ala Ser Glu Asn His Cys Tyr
```

1

```
<210> 111
<211> 9
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: TNF-ANTAGONIST
      PEPTIDE
<400> 111
Phe Cys Asn Ser Glu Asn His Cys Tyr
<210> 112
<211> 9
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: TNF-ANTAGONIST
      PEPTIDE
<400> 112
Phe Cys Asn Ser Glu Asn Arg Cys Tyr
<210> 113
<211> 10
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: TNF-ANTAGONIST
      PEPTIDE
<400> 113
Phe Cys Asn Ser Val Glu Asn Arg Cys Tyr
 1 5
```

```
<210> 114
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TNF-ANTAGONIST
      PEPTIDE
<400> 114
Tyr Cys Ser Gln Ser Val Ser Asn Asp Cys Phe
                 5
<210> 115
<211> 9
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TNF-ANTAGONIST
      PEPTIDE
<400> 115
Phe Cys Val Ser Asn Asp Arg Cys Tyr
                  5
<210> 116
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TNF-ANTAGONIST
      PEPTIDE
<400> 116
Tyr Cys Arg Lys Glu Leu Gly Gln Val Cys Tyr
                   5
```

<210> 117 ... < <211> 9 < <212> PRT

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<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TNF-ANTAGONIST
<400> 117
Tyr Cys Lys Glu Pro Gly Gln Cys Tyr
                5
<210> 118
<211> 9
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TNF-ANTAGONIST
<400> 118
Tyr Cys Arg Lys Glu Met Gly Cys Tyr
<210> 119
<211> 9
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TNF-ANTAGONIST
<400> 119
Phe Cys Arg Lys Glu Met Gly Cys Tyr
                  5
<210> 120
<211> 9
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TNF-ANTAGONIST
<400> 120
```

```
Tyr Cys Trp Ser Gln Asn Leu Cys Tyr
1 5
```

```
<210> 121
<211> 10
<212> PRT
<213> Artificial Sequence

<220>
<223> Description of Artificial Sequence:TNF-ANTAGONIST

<400> 121
Tyr Cys Glu Leu Ser Gln Tyr Leu Cys Tyr
1 5 10
```

```
<210> 122
<211> 9
<212> PRT
<213> Artificial Sequence

<220>
<223> Description of Artificial Sequence: TNF-ANTAGONIST

<400> 122
Tyr Cys Trp Ser Gln Asn Tyr Cys Tyr

1 5
```

```
<210> 123
<211> 9
<212> PRT
<213> Artificial Sequence

<220>
<223> Description of Artificial Sequence:TNF-ANTAGONIST

<400> 123
Tyr Cys Trp Ser Gln Tyr Leu Cys Tyr

1 5
```

<210> 124

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<211> 37

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EPO-MIMETIC PEPTIDE

<400> 124

5

Xaa Xaa Xaa Xaa Thr Trp Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa 25 20

Xaa Xaa Xaa Xaa Xaa 35

<210> 125

<211> 15

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: CTLA4-MIMETIC PEPTIDE

<400> 125

Gly Phe Val Cys Ser Gly Ile Phe Ala Val Gly Val Gly Arg Cys 10 5

<210> 126

<211> 15

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: CTLA4-MIMETIC PEPTIDE

<400> 126

Ala Pro Gly Val Arg Leu Gly Cys. Ala Val Leu Gly Arg Tyr Cys 10 5

```
<210> 127
<211> 27
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:C3B ANTAGONIST
<400> 127
Ile Cys Val Val Gln Asp Trp Gly His His Arg Cys Thr Ala Gly His
                                    10
Met Ala Asn Leu Thr Ser His Ala Ser Ala Ile
<210> 128
<211> 13
<212> PRT
<213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: C3B ANTAGONIST
       PEPTIDE
 <400> 128
 Ile Cys Val Val Gln Asp Trp Gly His His Arg Cys Thr
                  5
 <210> 129
 <211> 11
 <212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence: C3B ANTAGONIST
       PEPTIDE
  <400> 129
  Cys Val Val Gln Asp Trp Gly His His Ala Cys
        ··· 5
```

```
<210> 130
<211> 6
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:MDM/HDM
      ANTAGONIST PEPTIDE
<400> 130
Thr Phe Ser Asp Leu Trp
<210> 131
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:MDM/HDM
      ANTAGONIST PEPTIDE
<400> 131
Gln Glu Thr Phe Ser Asp Leu Trp Lys Leu Leu Pro
                  5
 <210> 132
 <211> 12
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:MDM/HDM
       ANTAGONIST PEPTIDE
 <400> 132
 Gln Pro Thr Phe Ser Asp Leu Trp Lys Leu Leu Pro
```

<210> 133 <211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:MDM/HDM
ANTAGONIST PEPTIDE

<400> 133

Gln Glu Thr Phe Ser Asp Tyr Trp Lys Leu Leu Pro 1 5 10

<210> 134

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:MDM/HDM ANTAGONIST PEPTIDE

<400> 134

Gln Pro Thr Phe Ser Asp Tyr Trp Lys Leu Leu Pro
1 5 10

<210> 135

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:MDM/HDM ANTAGONIST PEPTIDE

<400> 135

Met Pro Arg Phe Met Asp Tyr Trp Glu Gly Leu Asn 1 5 10

<210> 136

<211> 12

<212> PRT...

<213> Artificial Sequence

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<220> <223> Description of Artificial Sequence: C3B ANTAGONIST <400> 136 Val Gln Asn Phe Ile Asp Tyr Trp Thr Gln Gln Phe

<210> 137 <211> 12 <212> PRT <213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:MDM/HDM ANTAGONIST PEPTIDE

<400> 137 Thr Gly Pro Ala Phe Thr His Tyr Trp Ala Thr Phe 10

<210> 138 <211> 15 <212> PRT <213> Artificial Sequence

<220> <223> Description of Artificial Sequence:MDM/HDM ANTAGONIST PEPTIDE

<400> 138 Ile Asp Arg Ala Pro Thr Phe Arg Asp His Trp Phe Ala Leu Val 10 5

<210> 139 <211> 15 <212> PRT

<213> Artificial Sequence

<223> Description of Artificial Sequence:MDM/HDM ANTAGONIST PEPTIDE

```
<400> 139
Pro Arg Pro Ala Leu Val Phe Ala Asp Tyr Trp Glu Thr Leu Tyr
                      10 . 15
<210> 140
<211> 15
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:MDM/HDM
     ANTAGONIST PEPTIDE
<400> 140
Pro Ala Phe Ser Arg Phe Trp Ser Asp Leu Ser Ala Gly Ala His
                                 10
               5
<210> 141
<211> 15
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:MDM/HDM
     ANTAGONIST PEPTIDE
<400> 141
Pro Ala Phe Ser Arg Phe Trp Ser Lys Leu Ser Ala Gly Ala His
                                                    15
                                  10
       5
<210> 142
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:MDM/HDM
```

<400> 142 ...
Pro Xaa Phe Xaa Asp Tyr Trp Xaa Xaa Leu
1 5 10

ANTAGONIST PEPTIDE

```
<210> 143
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:MDM/HDM
     ANTAGONIST PEPTIDE
<400> 143
Gln Glu Thr Phe Ser Asp Leu Trp Lys Leu Leu Pro
               5
<210> 144
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:MDM/HDM
      ANTAGONIST PEPTIDE
<400> 144
Gln Pro Thr Phe Ser Asp Leu Trp Lys Leu Leu Pro
        5
<210> 145
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:MDM/HDM
      ANTAGONIST PEPTIDE
<400> 145
Gln Glu Thr Phe Ser Asp Tyr Trp Lys Leu Leu Pro
         5
```

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```
<210> 146
<211> 12
<212> PRT
```

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:MDM/HDM ANTAGONIST PEPTIDE

<400> 146 Gln Pro Thr Phe Ser Asp Tyr Trp Lys Leu Leu Pro 5

<210> 147 <211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SELECTIN ANTAGONIST PEPTIDE

<400> 147

Asp Ile Thr Trp Asp Gln Leu Trp Asp Leu Met Lys 5

<210> 148

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SELECTIN ANTAGONIST PEPTIDE

<400> 148

Asp Ile Thr Trp Asp Glu Leu Trp Lys Ile Met Asn 10 5 1

<210> 149 ...

<211> 12

<212> PRT

```
<213> Artificial Sequence
```

<220>

<223> Description of Artificial Sequence: SELECTIN ANTAGONIST PEPTIDE

<400> 149

Asp Tyr Thr Trp Phe Glu Leu Trp Asp Met Met Gln
1 5 10

<210> 150

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SELECTIN ANTAGONIST PEPTIDE

<400> 150

Gln Ile Thr Trp Ala Gln Leu Trp Asn Met Met Lys
1 5 10

<210> 151

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:MDM/HDM ANTAGONIST PEPTIDE

<400> 151

Asp Met Thr Trp His Asp Leu Trp Thr Leu Met Ser

1 5 10

<210> 152

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:MDM/HDM ANTAGONIST PEPTIDE

<400> 152 Asp Tyr Ser Trp His Asp Leu Trp Glu Met Met Ser 1 5 10

<210> 153

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:MDM/HDM ANTAGONIST PEPTIDE

<400> 153

Glu Ile Thr Trp Asp Gln Leu Trp Glu Val Met Asn
1 5 10

<210> 154

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:MDM/HDM ANTAGONIST PEPTIDE

<400> 154

His Val Ser Trp Glu Gln Leu Trp Asp Ile Met Asn 1 5 10

<210> 155

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SELECTIN ANTAGONIST PEPTIDE

<210> 156

<211> 13

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SELECTIN ANTAGONIST PEPTIDE

<400> 156

Arg Asn Met Ser Trp Leu Glu Leu Trp Glu His Met Lys
1 5 10

<210> 157

<211> 18

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SELECTIN

<400> 157

Ala Glu Trp Thr Trp Asp Gln Leu Trp His Val Met Asn Pro Ala Glu

1 5 10 15

Ser Gln

<210> 158

<211> 14

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SELECTIN

<400> 158

His Arg Ala Glu Trp Leu Ala Leu Trp Glu Gln Met Ser Pro

1 5 10

<210> 159

<211> 14

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SELECTIN ANTAGONIST PEPTIDE

<400> 159

Lys Lys Glu Asp Trp Leu Ala Leu Trp Arg Ile Met Ser Val

<210> 160

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SELECTIN

<400> 160

Ile Thr Trp Asp Gln Leu Trp Asp Leu Met Lys
1 5 10

<210> 161

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SELECTIN

<400> 161

Asp Ile Thr Trp Asp Gln Leu Trp Asp Leu Met Lys
1 5 10

<210> 162

```
<211> 12
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: SELECTIN
<400> 162
Asp Ile Thr Trp Asp Gln Leu Trp Asp Leu Met Lys
                  5
<210> 163
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SELECTIN
     ANTAGONIST PEPTIDE
<400> 163
Asp Ile Thr Trp Asp Gln Leu Trp Asp Leu Met Lys
                  5
<210> 164
<211> 13
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: CALMODULIN
      ANTAGONIST PEPTIDE
<400> 164
Ser Cys Val Lys Trp Gly Lys Lys Glu Phe Cys Gly Ser
  1
<210> 165
<211> 12
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<212> PRT ...

<213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: CALMODULIN
<400> 165
Ser Cys Trp Lys Tyr Trp Gly Lys Glu Cys Gly Ser
                 5
<210> 166
<211> 13
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:CALMODULIN
      ANTAGONIST PEPTIDE
<400> 166
Ser Cys Tyr Glu Trp Gly Lys Leu Arg Trp Cys Gly Ser
                                     10
                 5
<210> 167
<211> 13
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: CALMODULIN
      ANTAGONIST PEPTIDE
<400> 167
Ser Cys Leu Arg Trp Gly Lys Trp Ser Asn Cys Gly Ser
<210> 168
 <211> 13
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: CALMODULIN
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ANTAGONIST PEPTIDE

```
<400> 168
Ser Cys Trp Arg Trp Gly Lys Tyr Gln Ile Cys Gly Ser
                  5
<210> 169
<211> 13
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: CALMODULIN
      ANTAGONIST PEPTIDE
<400> 169
Ser Cys Val Ser Trp Gly Ala Leu Lys Leu Cys Gly Ser
                  5
  1
<210> 170
<211> 13
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: CALMODULIN
      ANTAGONIST PEPTIDE
<400> 170
Ser Cys Ile Arg Trp Gly Gln Asn Thr Phe Cys Gly Ser
                  5
<210> 171
<211> 13
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: CALMODULIN
      ANTAGONIST PEPTIDE
<400> 171
Ser Cys Trp Gln Trp Gly Asn Leu Lys Ile Cys Gly Ser
```

10

5 .

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<210> 172
<211> 13
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:CALMODULIN
      ANTAGONIST PEPTIDE
<400> 172
Ser Cys Val Arg Trp Gly Gln Leu Ser Ile Cys Gly Ser
                 5
<210> 173
<211> 21
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: CALMODULIN
      ANTAGONIST PEPTIDE
<400> 173
Leu Lys Lys Phe Asn Ala Arg Arg Lys Leu Lys Gly Ala Ile Leu Thr
                                     10
                 5
Thr Met Leu Ala Lys
             20
<210> 174
<211> 18
 <212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence:CALMODULIN
 Arg Arg Trp Lys Lys Asn Phe Ile Ala Val Ser Ala Ala Asn Arg Phe
                                                          15
                                      10
                  5
```

Lys Lys

<210> 175

<211> 18

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:CALMODULIN

<400> 175

Arg Lys Trp Gln Lys Thr Gly His Ala Val Arg Ala Ile Gly Arg Leu
1 5 10 15

Ser Ser

<210> 176

<211> 14

<212> PRT

<213> Artificial Sequence '

<220>

<223> Description of Artificial Sequence:CALMODULIN
 ANTAGONIST PEPTIDE

<400> 176

Ile Asn Leu Lys Ala Leu Ala Ala Leu Ala Lys Lys Ile Leu
1 5 10

<210> 177

<211> 18

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:CALMODULIN ANTAGONIST PEPTIDE

<400> 177

Lys Ile Trp Ser Ile Leu Ala Pro Leu Gly Thr Thr Leu Val Lys Leu

1 5 10 15

Val Ala

<210> 178

<211> 14

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:CALMODULIN ANTAGONIST PEPTIDE

<400> 178

Leu Lys Lys Leu Leu Lys Leu Lys Lys Leu Leu Lys Leu 1 5 10

<210> 179

<211> 18

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:CALMODULIN ANTAGONIST PEPTIDE

<400> 179

Leu Lys Trp Lys Lys Leu Leu Lys Leu Leu Lys Lys Leu Leu Lys Lys

1 5 10 15

Leu Leu

<210> 180

<211> 17

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:CALMODULIN ANTAGONIST PEPTIDE

<400> 180 Ala Glu Trp Pro Ser Leu Thr Glu Ile Lys Thr Leu Ser His Phe Ser 10 Val <210> 181 <211> 17 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: CALMODULIN ANTAGONIST PEPTIDE <400> 181 Ala Glu Trp Pro Ser Pro Thr Arg Val Ile Ser Thr Thr Tyr Phe Gly 10 5 Ser <210> 182 <211> 17 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: CALMODULIN ANTAGONIST PEPTIDE <400> 182 Ala Glu Leu Ala His Trp Pro Pro Val Lys Thr Val Leu Arg Ser Phe 5 10 Thr

<210> 183 <211> 17

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<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:CALMODULIN
   ANTAGONIST PEPTIDE
<400> 183
Ala Glu Gly Ser Trp Leu Gln Leu Leu Asn Leu Met Lys Gln Met Asn
                  5
                                     10
Asn
<210> 184
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:CALMODULIN
      ANTAGONIST PEPTIDE
<400> 184
Ala Glu Trp Pro Ser Leu Thr Glu Ile Lys
                  5
  1
<210> 185
<211> 27
<212> PRT
<213> Artificial Sequence
<220>
 <223> Description of Artificial
       Sequence: VINCULIN-BINDING PEPTIDE
 <400> 185
 Ser Thr Gly Gly Phe Asp Asp Val Tyr Asp Trp Ala Arg Gly Val Ser
                                     10
                   5
```

· 25

Ser Ala Leu Thr Thr Thr Leu Val Ala Thr Arg

··· 20

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WO 00/24782 <210> 186 <211> 27 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: VINCULIN-BINDING PEPTIDE <400> 186 Ser Thr Gly Gly Phe Asp Asp Val Tyr Asp Trp Ala Arg Arg Val Ser 5 -Ser Ala Leu Thr Thr Thr Leu Val Ala Thr Arg 20 <210> 187 <211> 30 <212> PRT

<213> Artificial Sequence

<223> Description of Artificial Sequence: VINCULIN BINDING PEPTIDE

<400> 187

Ser Arg Gly Val Asn Phe Ser Glu Trp Leu Tyr Asp Met Ser Ala Ala 10 15

Met Lys Glu Ala Ser Asn Val Phe Pro Ser Arg Arg Ser Arg 20 . 25

<210> 188

<211> 30

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: VINCULIN BINDING PEPTIDE

<400> 188

Ser Ser Gln Asn Trp Asp Met Glu Ala Gly Val Glu Asp Leu Thr Ala

1 5 10 15

Ala Met Leu Gly Leu Leu Ser Thr Ile His Ser Ser Ser Arg
20 25 30

<210> 189

<211> 31

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VINCULIN BINDING PEPTIDE

<400> 189

Ser Ser Pro Ser Leu Tyr Thr Gln Phe Leu Val Asn Tyr Glu Ser Ala 1 5 10 15

Ala Thr Arg Ile Gln Asp Leu Leu Ile Ala Ser Arg Pro Ser Arg 20 25 30

<210> 190

<211> 31

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VINCULIN BINDING PEPTIDE

<400> 190

Ser Ser Thr Gly Trp Val Asp Leu Leu Gly Ala Leu Gln Arg Ala Ala 1 5 10 15

Asp Ala Thr Arg Thr Ser Ile Pro Pro Ser Leu Gln Asn Ser Arg

<210> 191

<211> 18

<212> PRT ...

<213> Artificial Sequence

```
<220>
<223> Description of Artificial Sequence: VINCULIN
      BINDING PEPTIDE
<400> 191
Asp Val Tyr Thr Lys Lys Glu Leu Ile Glu Cys Ala Arg Arg Val Ser
                                   10
Glu Lys
<210> 192
<211> 22
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:C4BP-BINDING
      PEPTIDE
<400> 192
Glu Lys Gly Ser Tyr Tyr Pro Gly Ser Gly Ile Ala Gln Phe His Ile
                                    10
                 5
Asp Tyr Asn Asn Val Ser
             20
<210> 193
<211> 22
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:C4BP-BINDING
      PEPTIDE
<400> 193
Ser Gly Ile Ala Gln Phe His Ile Asp Tyr Asn Asn Val Ser Ser Ala
                                     10
Glu Gly Trp His Val Asn
```

20

```
<210> 194
 <211> 34
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:C4BP-BINDING
       PEPTIDE
 <400> 194
 Leu Val Thr Val Glu Lys Gly Ser Tyr Tyr Pro Gly Ser Gly Ile Ala
                   5
 Gln Phe His Ile Asp Tyr Asn Asn Val Ser Ser Ala Glu Gly Trp His
                                  25
              20
Val Asn
 <210> 195
 <211> 14
 <212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence:C4BP-BINDING
       PEPTIDE
 <400> 195
 Ser Gly Ile Ala Gln Phe His Ile Asp Tyr Asn Asn Val Ser
                   5
  <210> 196
  <211> 17
  <212> PRT
  <213> Artificial Sequence
  <223> Description of Artificial Sequence: UKR ANTAGONIST
        PEPTIDE
```

Ala Glu Pro Met Pro His Ser Leu Asn Phe Ser Gln Tyr Leu Trp Tyr

<400> 196

1 5 10 15

Thr

<210> 197

<211> 17

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:UKR ANTAGONIST
 PEPTIDE

<400> 197

Ala Glu His Thr Tyr Ser Ser Leu Trp Asp Thr Tyr Ser Pro Leu Ala 1 5 10 15

Phe

<210> 198

<211> 17

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial
 Sequence:VINCULIN-BINDING PEPTIDE

<400> 198

Ala Glu Leu Asp Leu Trp Met Arg His Tyr Pro Leu Ser Phe Ser Asn 1 5 10 15

Arg

<210> 199

<211> 17

<212> PRT ...

<213> Artificial Sequence

```
<220>
<223> Description of Artificial Sequence: UKR ANTAGONIST
     PEPTIDE
<400> 199
Ala Glu Ser Ser Leu Trp Thr Arg Tyr Ala Trp Pro Ser Met Pro Ser
                                    10
                 5
Tyr
<210> 200
<211> 17
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:UKR ANTAGONIST
      PEPTIDE
<400> 200
Ala Glu Trp His Pro Gly Leu Ser Phe Gly Ser Tyr Leu Trp Ser Lys
                                                         15
                  5
                                     10
Thr
<210> 201
<211> 17
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:UKR ANTAGONIST
      PEPTIDE
<400> 201
Ala Glu Pro Ala Leu Leu Asn Trp Ser Phe Phe Asn Pro Gly Leu
                                    10
                 5
```

His

```
<210> 202
<211> 17
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:UKR ANTAGONIST
      PEPTIDE
<400> 202
Ala Glu Trp Ser Phe Tyr Asn Leu His Leu Pro Glu Pro Gln Thr Ile
                 5
                                  10
Phe
<210> 203
<211> 17
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:UKR ANTAGONIST
      PEPTIDE
Ala Glu Pro Leu Asp Leu Trp Ser Leu Tyr Ser Leu Pro Pro Leu Ala
                                   10
                                                      15
                  5
Met
 <210> 204
 <211> 17
 <212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence: UKR ANTAGONIST
      PEPTIDE
 Ala Glu Pro Thr Leu Trp Gln Leu Tyr Gln Phe Pro Leu Arg Leu Ser
```

1 5 10 15

Gly

<210> 205

<211> 17

<212> PRT

<213> Artificial Sequence

<220>

<400> 205

Ala Glu Ile Ser Phe Ser Glu Leu Met Trp Leu Arg Ser Thr Pro Ala 1 5 10 15

Phe

<210> 206

<211> 17

<212> PRT

<213> Artificial Sequence

<220>

<400> 206

Ala Glu Leu Ser Glu Ala Asp Leu Trp Thr Thr Trp Phe Gly Met Gly

1 5 10 15

Ser

<210> 207

<211> 17

<212> PRT-

<213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: UKR ANTAGONIST
      PEPTIDE
<400> 207
Ala Glu Ser Ser Leu Trp Arg Ile Phe Ser Pro Ser Ala Leu Met Met
                                   10
                 5
Ser
<210> 208
<211> 17
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:UKR ANTAGONIST
      PEPTIDE
<400> 208
Ala Glu Ser Leu Pro Thr Leu Thr Ser Ile Leu Trp Gly Lys Glu Ser
                                    10
                  5
Val
<210> 209
<211> 17
<212> PRT
<213> Artificial Sequence
 <223> Description of Artificial Sequence: UKR ANTAGONIST
      PEPTIDE
```

Ala Glu Thr Leu Phe Met Asp Leu Trp His Asp Lys His Ile Leu Leu

5

<400> 209

Thr

```
<210> 210
<211> 17
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: UKR ANTAGONIST
      PEPTIDE
<400> 210
Ala Glu Ile Leu Asn Phe Pro Leu Trp His Glu Pro Leu Trp Ser Thr
                                    10
                  5
Glu
<210> 211
<211> 17
<212> PRT
<213> Artificial Sequence
 <223> Description of Artificial Sequence:UKR ANTAGONIST
      PEPTIDE
 <400> 211
Ala Glu Ser Gln Thr Gly Thr Leu Asn Thr Leu Phe Trp Asn Thr Leu
                                                         15
                                     10
 Arg
 <210> 212
 <211> 9
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:IL-1 ANTAGONIST
       PEPTIDE
 <220>
 <223> At position 1, Xaa is V, L, I, E, P, G, Y, M, T,
```

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or D

<220>

<223> At position 2, Xaa is Y, W or F

<220>

<223> At position 3, Xaa is E, F, V, W or Y

<220>

<223> At position 5, Xaa is P or azetidine

<220>

<223> At position 7, Xaa is S, A, V or L

<223> At position 8, Xaa is M, F, V, R, Q, K, T, S, D, L, I or E

<220>

<223> At position 9, Xaa is E, L, W, V, H, I, G, A, D, L, Y, N, Q or P

<400> 212

Xaa Xaa Xaa Gln Xaa Tyr Xaa Xaa Xaa 1 5

<210> 213

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 213

Thr Ala Asn Val Ser Ser Phe Glu Trp Thr Pro Tyr Tyr Trp Gln Pro 10 1

Tyr Ala Leu Pro Leu

20

<210> 214

<211> 18

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```
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 214
Ser Trp Thr Asp Tyr Gly Tyr Trp Gln Pro Tyr Ala Leu Pro Ile Ser
                                    10
                 5
Gly Leu
<210> 215
<211> 21
<212> PRT
<213> Artificial Sequence
<220>
 <223> Description of Artificial Sequence:IL-1 ANTAGONIST
       PEPTIDE
 <400> 215
 Glu Thr Pro Phe Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro
                                                         15
                                     10
 Tyr Ala Leu Pro Leu
             20
 <210> 216
 <211> 21
 <212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
```

PEPTIDE

<400> 216

Glu Asn Thr Tyr Ser Pro Asn Trp Ala Asp Ser Met Tyr Trp Gln Pro ... 5

Tyr Ala Leu Pro Leu

20

```
<210> 217
<211> 21
<212> PRT
<213> Artificial Sequence
<220>
```

<400> 217
Ser Val Gly Glu Asp His Asn Phe Trp Thr Ser Glu Tyr Trp Gln Pro
1 5 10 15

Tyr Ala Leu Pro Leu 20

<210> 218 <211> 21 <212> PRT <213> Artificial Sequence

<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

Tyr Ala Leu Pro Leu 20

<210> 219 <211> 11 <212> PRT <213> Artificial Sequence

<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST
PEPTIDE

```
<400> 219
Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro Tyr
 1 5
<210> 220
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<400> 220
Phe Glu Trp Thr Pro Gly Tyr Trp Gln His Tyr
<210> 221
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<220>
<223> At position 10, Xaa=azetidine
<400> 221
Phe Glu Trp Thr Pro Gly Trp Tyr Gln Xaa Tyr
                 5
<210> 222
<211> 11
<212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
```

```
<220>
  <223> At position 1, optionally acetylated at N-terminus
  <220>
<223> At position 10, Xaa=azetidine
  <400> 222
  Phe Glu Trp Thr Pro Gly Trp Tyr Gln Xaa Tyr
                   5
  <210> 223
  <211> 12
  <212> PRT
  <213> Artificial Sequence
  <220>
  <223> Description of Artificial Sequence: IL-1 ANTAGONIST
        PEPTIDE
  <220>
  <223> At position 11, Xaa=azetidine
  <400> 223
  Phe Glu Trp Thr Pro Gly Trp Pro Tyr Gln Xaa Tyr
                                       10
                    5
  <210> 224
  <211> 11
  <212> PRT
  <213> Artificial Sequence
  <220>
  <223> Description of Artificial Sequence: IL-1 ANTAGONIST
         PEPTIDE
   <220>
   <223> At position 10, Xaa=azetidine
   <400> 224
   Phe Ala Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr
                    5
```

```
<210> 225
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<220>
<223> At position 10, Xaa=azetidine
<400> 225
Phe Glu Trp Ala Pro Gly Tyr Trp Gln Xaa Tyr
      5
<210> 226
<211> 11
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 10, Xaa=azetidine
<400> 226
Phe Glu Trp Val Pro Gly Tyr Trp Gln Xaa Tyr
                  5
 <210> 227
 <211> 11
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <220>
<223> At position 10, Xaa=azetidine
```

<400> 227

```
Phe Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr
                 5
<210> 228
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST
     PEPTIDE
<220>
<223> At position 1, optionally acetylated at N-terminus
<223> At position 10, Xaa=azetidine
<400> 228
Phe Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr
                5
<210> 229
<211> 11
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<220>
<223> At position 6, products="MeGly"
<220>
<223> At position 10, Xaa=azetidine
<400> 229
Phe Glu Trp Thr Pro Xaa Trp Tyr Gln Xaa Tyr
 1 ... 5 . 10
```

```
<210> 230
<211> 11
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 6, Xaa=MeGly
<220>
<223> At position 10, Xaa=azetidine
<400> 230
Phe Glu Trp Thr Pro Xaa Trp Tyr Gln Xaa Tyr
                 5
<210> 231
<211> 11
<212> PRT
<213> Artificial Sequence
<220> ·
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 231
Phe Glu Trp Thr Pro Gly Tyr Tyr Gln Pro Tyr
                                     10.
                 5
<210> 232
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST
      PEPTIDE
```

Phe Glu Trp Thr Pro Gly Trp Trp Gln Pro Tyr

1 5 10

<210> 233

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 233

Phe Glu Trp Thr Pro Asn Tyr Trp Gln Pro Tyr
1 5 10

<210> 234

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<220>

<223> At position 5, Xaa=pipecolic acid

<220>

<223> At position 10, Xaa=azetidine

<400> 234

Phe Glu Trp Thr Xaa Val Tyr Trp Gln Xaa Tyr
1 5 10

<210> 235

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

```
<220>
<223> At position 5, Xaa=pipecolic acid
<220>
<223> At position 10, Xaa=azetidine
<400> 235
Phe Glu Trp Thr Xaa Gly Tyr Trp Gln Xaa Tyr
                 5
<210> 236
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 6, Xaa=Aib
<220>
<223> At position 10, Xaa=azetidine
<400> 236
Phe Glu Trp Thr Pro Xaa Tyr Trp Gln Xaa Tyr
                  5
<210> 237
<211> 11
<212> PRT
<213> Artificial Sequence
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <220>
 <223> At position 5, Xaa=MeGly
 <220>
 <223> At position 10, Xaa=azetidine
```

```
<400> 237
Phe Glu Trp Thr Xaa Gly Tyr Trp Gln Xaa Tyr
<210> 238
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
 <223> At position 11, amino group added at C-terminus
<400> 238.
 Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro Tyr
                   5
                                      10
  1
 <210> 239
 <211> 11
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <223> At position 11, amino group added at C-terminus
 <400> 239
 Phe Glu Trp Thr Pro Gly Tyr Trp Gln His Tyr
                   5
 <210> 240
 <211> 11
<212> PRT "
```

<213> Artificial Sequence

```
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 10, Xaa is an azetidine residue
<220>
<223> At position 11 amino group added at C-terminus
<400> 240
Phe Glu Trp Thr Pro Gly Trp Tyr Gln Xaa Tyr
                 5
<210> 241
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 1, optionally acetylated at
      N-terminus
<220>
<223> At position 10, Xaa is an azetidine residue
<220>
<223> At position 11 amino group added at C-terminus
Phe Glu Trp Thr Pro Gly Trp Tyr Gln Xaa Tyr
                 5
<210> 242
<211> 11
<212> PRT
 <213> Artificial Sequence
<220>
```

<223> Description of Artificial Sequence: IL-1 ANTAGONIST

PEPTIDE

<220>

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<220>
<223> At position 8, Xaa is a phyosphotyrosyl residue
<220>
<223> At position 10, Xaa is an azetidine residue
<220>
<223> At position 11, amino group added at C-terminus
<400> 242
Phe Glu Trp Thr Pro Gly Trp Xaa Gln Xaa Tyr
                  5
<210> 243
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<223> At position 10, Xaa is an azetidine residue
<223> At position 11 amino group added at C-terminus
<400> 243
Phe Ala Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr
                  5
<210> 244
<211> 11
```

```
<211> 11
<212> PRT
<213> Artificial Sequence

<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE
```

```
<223> At position 10, Kaa is an azetidine residue
<220>
<223> At position 11 amino group added at C-terminus
<400> 244
Phe Glu Trp Ala Pro Gly Tyr Trp Gln Xaa Tyr
                  5
<210> 245
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 10, Xaa is an azetidine residue
<220>
<223> At position 11 amino group added at C-terminus
<400> 245
Phe Glu Trp Val Pro Gly Tyr Trp Gln Xaa Tyr
                  5
  1
<210> 246
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 10, Xaa is an azetidine residue
<223> At position 11 amino group added at C-terminus
<400> 246
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Phe Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr 1 5 10

<210> 247

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<220>

<223> At position 1 acetylated at N-terminus

<220>

<223> At position 10, Xaa is an azetidine residue

<220>

<223> At position 11 amino group added at C-terminus

<400> 247

Xaa Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr
1 5 10

<210> 248

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<220>

<223> At position 6, D amino acid residue

<220>

<223> At position 10, Xaa is an azetidine residue

<220>

<223> At position 11 amino group added at C-terminus

<400> 248

Phe Glu Trp Thr Pro Ala Trp Tyr Gln Xaa Tyr
1 5 10

<210> 249

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<220>

<223> At position 6, Xaa is a sarcosine residue

2220>

<223> At position 10, Xaa is an azetidine residue

<220>

<223> At position 11 amino group added at C-terminus

<400> 249

Phe Glu Trp Thr Pro Xaa Trp Tyr Gln Xaa Tyr
1 5 10

<210> 250

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<220>

<223> At position 11 amino group added at C-terminus

<400> 250

Phe Glu Trp Thr Pro Gly Tyr Tyr Gln Pro Tyr

1 5 10

<210> 251

```
<211> 11
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:IL-1 ANTAGONIST
     PEPTIDE
<220>
<223> At position 11 amino group added at C-terminus
<400> 251
Phe Glu Trp Thr Pro Gly Trp Trp Gln Pro Tyr
    · 5
                                    10
<210> 252
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<223> At position 11 amino group added at C-terminus
Phe Glu Trp Thr Pro Asn Tyr Trp Gln Pro Tyr
                5
<210> 253
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <220>
 <223> At position 6, D amino acid residue
```

<220>

```
<223> At position 10, Xaa is an azetidine residue
<223> At position 11, amino group added at C-terminus
<400> 253
Phe Glu Trp Thr Pro Val Tyr Trp Gln Xaa Tyr
                  5
<210> 254
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 5, Xaa is a pipecolic acid residue
<220>
<223> At position 10, Xaa is an azetidine residue
<220>
<223> At position 11, amino group added at C-terminus
<400> 254
Phe Glu Trp Thr Xaa Gly Tyr Trp Gln Xaa Tyr
  1
 <210> 255
 <211> 11
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <223> At position 6, Xaa=pipecolic acid
 <220>
```

```
<223> At position 10, Xaa=azetidine
<400> 255
Phe Glu Trp Thr Pro Xaa Tyr Trp Gln Xaa Tyr
                 5
<210> 256
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 5, Xaa=MeGly
<220>
<223> At position 10, Xaa=azetidine
<400> 256
Phe Glu Trp Thr Xaa Gly Tyr Trp Gln Xaa Tyr
<210> 257
<211> 15
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: INTEGRIN
      BINDING PEPTIDE
Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro Tyr Ala Leu Pro Leu
                                   10
  1
                  5
<210> 258
<211> 11 ...
<212> PRT
```

<213> Artificial Sequence

```
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 1, Xaa is a 1-naphthylalanine residue
<220>
<223> At position 10, Xaa is an azetidine residue
<220>
<223> At position 11, amino group added at C-terminus
<400> 258
Xaa Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Tyr
                  5
<210> 259
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 10, Xaa is a azetidine residue
<223> At position 11, amino group added at C-terminus
<400> 259
Tyr Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Tyr
                   5
  1
 <210> 260
 <211> 11
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
```

PEPTIDE

```
<220>
<223> At position 10, Xaa is an azetidine residue

<220>
<223> At position 11, amino group added at C-terminus

<400> 260

Phe Glu Trp Val Pro Gly Tyr Tyr Gln Xaa Tyr

1 5 10
```

<210> 261 <211> 11 <212> PRT <213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<220>

<223> At position 6, D amino acid residue

<220>

<223> At position 10, Xaa is an azetidine residue

<220>

<223> At position 11, amino group added at C-terminus

<400> 261

Phe Glu Trp Thr Pro Ser Tyr Tyr Gln Xaa Tyr 1 5 10

<210> 262

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<220>

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<223> At position 6, D amino acid residue
<223> At position 10, Xaa is an azetidine residue
<220>
<223> At position 11, amino group added at C-terminus
<400> 262
Phe Glu Trp Thr Pro Asn Tyr Tyr Gln Xaa Tyr
<210> 263
<211> 4
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST
      PEPTIDE
<400> 263
Thr Lys Pro Arg
<210> 264
<211> 5
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 264
Arg Lys Ser Ser Lys
 <210> 265
 <211> 5 "
 <212> PRT
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<213> Artificial Sequence

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<220>
  <223> Description of Artificial Sequence: IL-1 ANTAGONIST
        PEPTIDE
  <400> 265
  Arg Lys Gln Asp Lys
  <210> 266
  <211> 6
  <212> PRT
  <213> Artificial Sequence
<220>
  <223> Description of Artificial Sequence: IL-1 ANTAGONIST
        PEPTIDE
  <400> 266
  Asn Arg Lys Gln Asp Lys
  <210> 267
  <211> 6
  <212> PRT
  <213> Artificial Sequence
  <220>
  <223> Description of Artificial Sequence: IL-1 ANTAGONIST
        PEPTIDE
  <400> 267
  Arg Lys Gln Asp Lys Arg
   1
  <210> 268
  <211> 9
  <212> PRT
  <213> Artificial Sequence
  <220>
  <223> Description of Artificial Sequence: IL-1 ANTAGONIST
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PEPTIDE

```
<400> 268
Glu Asn Arg Lys Gln Asp Lys Arg Phe
               5
```

<210> 269 <211> 6 <212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 269 Val Thr Lys Phe Tyr Phe 5

<210> 270

<211> 5

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 270

Val Thr Lys Phe Tyr 1

<210> 271

<211> 5

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 271

```
Val Thr Asp Phe Tyr
<210> 272
<211> 17
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<400> 272
Ser Gly Ser Gly Val Leu Lys Arg Pro Leu Pro Ile Leu Pro Val Thr
                                 10 15
                 5
Arg
<210> 273
<211> 17
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:MCA/MCP
      PROTEASE INHIBITOR PEPTIDE
<400> 273
Arg Trp Leu Ser Ser Arg Pro Leu Pro Pro Leu Pro Leu Pro Pro Arg
                                  10
          5
Thr
<210> 274
<211> 20
<212> PRT
<213> Artificial Sequence
```

<223> Description of Artificial Sequence: MCA/MCPPROTEASE

<220>

INHIBITOR PEPTIDE

<400> 274

Gly Ser Gly Ser Tyr Asp Thr Leu Ala Leu Pro Ser Leu Pro Leu His

1 5 10 15

Pro Met Ser Ser

20

<210> 275

<211> 20

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:MCA/MCP
 PROTEASE INHIBITOR PEPTIDE

<400> 275

Gly Ser Gly Ser Tyr Asp Thr Arg Ala Leu Pro Ser Leu Pro Leu His 1 5 10 15

Pro Met Ser Ser 20

<210> 276

<211> 20

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:MCA/MCP
 PROTEASE INHIBITOR PEPTIDE

<400> 276

Gly Ser Gly Ser Ser Gly Val Thr Met Tyr Pro Lys Leu Pro Pro His 1 5 10 15

Trp Ser Met Ala

20

<210> 277

```
<211> 20
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:MCA/MCP
      PROTEASE INHIBITOR PEPTIDE
<400> 277
Gly Ser Gly Ser Ser Gly Val Arg Met Tyr Pro Lys Leu Pro Pro His
                                    10
Trp Ser Met Ala
        20
<210> 278
<211> 20
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:MCA/MCP
      PROTEASE INHIBITOR PEPTIDE
<400> 278
Gly Ser Gly Ser Ser Ser Met Arg Met Val Pro Thr Ile Pro Gly Ser
                  5
                                     10
 1
Ala Lys His Gly
             20
·<210> 279
<211> 6
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: ANTI-HBV
      PEPTIDE
 <400> 279
Leu Leu Gly Arg Met Lys
  1
                 5
```

```
<210> 280
<211> 8
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:ANTI-HBV
      PEPTIDE
<400> 280
Ala Leu Leu Gly Arg Met Lys Gly
<210> 281
<211> 6
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: ANTI-HBV
      PEPTIDE
<400> 281
Leu Asp Pro Ala Phe Arg
 1
<210> 282
<211> 7
<212> PRT
 <213> Artificial Sequence
<223> Description of Artificial Sequence: SH3 ANTAGONIST
 <400> 282
 Arg Pro Leu Pro Pro Leu Pro
```

<210> 283 <211> 7

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<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 283
Arg Glu Leu Pro Pro Leu Pro
                5
<210> 284
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: MSH3 ANTAGONIST
<400> 284
Ser Pro Leu Pro Pro Leu Pro
          5
<210> 285
<211> 7
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 285
Gly Pro Leu Pro Pro Leu Pro
          5
 1
<210> 286
<211> 7
<212> PRT
<213> Artificial Sequence
<220> ...
<223> Description of Artificial Sequence: SH3 ANTAGONIST
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<400> 286
Arg Pro Leu Pro Ile Pro Pro
                5
<210> 287
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:MAST CELL
      ANTAGONISTS/MAST CELL PROTEASE INHIBITOR
<400> 287
Arg Pro Leu Pro Ile Pro Pro
                  5
 1
<210> 288
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 288
Arg Arg Leu Pro Pro Thr Pro
                5
 1
<210> 289
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 289
Arg Gln Leu Pro Pro Thr Pro
      ··· 5
```

```
<210> 290
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 290
Arg Pro Leu Pro Ser Arg Pro
  1
<210> 291
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 291
Arg Pro Leu Pro Thr Arg Pro
<210> 292
<211> 7
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 292
Ser Arg Leu Pro Pro Leu Pro
                   5
 <210> 293
 <211> 7
 <212> PRT
 <213> Artificial Sequence
```

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<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 293
Arg Ala Leu Pro Ser Pro Pro
         5
<210> 294
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 294
Arg Arg Leu Pro Arg Thr Pro
<210> 295
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 295
Arg Pro Val Pro Pro Ile Thr
<210> 296
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 296...
Ile Leu Ala Pro Pro Val Pro
         5
```

```
<210> 297
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 297
Arg Pro Leu Pro Met Leu Pro
  1
                5
<210> 298
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 298
Arg Pro Leu Pro Ile Leu Pro
<210> 299
<211> 7
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 299
Arg Pro Leu Pro Ser Leu Pro
                 5
  1
```

<210> 300 ···· <211> 7 <212> PRT

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<213> Artificial Sequence
  <220>
  <223> Description of Artificial Sequence: SH3 ANTAGONIST
  <400> 300
  Arg Pro Leu Pro Ser Leu Pro
  <210> 301
  <211> 7
  <212> PRT
  <213> Artificial Sequence
  <223> Description of Artificial Sequence:SH3 ANTAGONIST
  <400> 301
  Arg Pro Leu Pro Met Ile Pro
   1
  <210> 302
  <211> 7
  <212> PRT
  <213> Artificial Sequence
  <220>
  <223> Description of Artificial Sequence: SH3 ANTAGONIST
  <400> 302
  Arg Pro Leu Pro Leu Ile Pro
                   5
  <210> 303
  <211> 7
  <212> PRT
<213> Artificial Sequence
  <223> Description of Artificial Sequence: SH3 ANTAGONIST
  <400> 303
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```
Arg Pro Leu Pro Pro Thr Pro
 1
<210> 304
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
.<400> 304
Arg Ser Leu Pro Pro Leu Pro
<210> 305
<211> 7
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 305
Arg Pro Gln Pro Pro Pro Pro
                 5
<210> 306
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 306
Arg Gln Leu Pro Ile Pro Pro
  1
```

<210> 307

```
<211> 12
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 307
Xaa Xaa Xaa Arg Pro Leu Pro Pro Leu Pro Xaa Pro
                  5
<210> 308
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 308
Xaa Xaa Xaa Arg Pro Leu Pro Pro Ile Pro Xaa Xaa
                  5
<210> 309
<211> 12
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 309
Xaa Xaa Xaa Arg Pro Leu Pro Pro Leu Pro Xaa Xaa
                                     10
                 5
<210> 310
 <211> 12
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: SH3 ANTAGONIST
```

```
<400> 310
Arg Xaa Xaa Arg Pro Leu Pro Pro Leu Pro Xaa Pro
       5
<210> 311
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 311
Arg Xaa Xaa Arg Pro Leu Pro Pro Leu Pro Pro Pro
                  5
<210> 312
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 312
Pro Pro Pro Tyr Pro Pro Pro Pro Ile Pro Xaa Xaa
                                    10
                 5
<210> 313
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 313
 Pro Pro Pro Tyr Pro Pro Pro Pro Val Pro Xaa Xaa
```

5

```
<210> 314
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 314
Leu Xaa Xaa Arg Pro Leu Pro Xaa Xaa Pro
                  5
<210> 315
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<220>
<223> At position 1, Xaa is an aliphatic amino acid
      residue
<400> 315
Xaa Xaa Xaa Arg Pro Leu Pro Xaa Leu Pro
                 5
<210> 316
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<220>
<223> At position 4, Xaa is an aromatic amino acid
      residue
<220>
<223> At position 9, Xaa is an aliphatic amino acid
```

residue

```
<400> 316
Pro Pro Xaa Xaa Tyr Pro Pro Pro Xaa Pro
1 5 10
```

<210> 317

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SH3 ANTAGONIST

<220>

<223> At position 1, Xaa is a basic amino acid residue

<220>

<223> At position 4, Xaa is an aliphatic amino acid residue

<400> 317

Xaa Pro Pro Xaa Pro Xaa Lys Pro Xaa Trp Leu 1 5 10

<210> 318

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SH3 ANTAGONIST

<220>

<223> At position 4, Xaa is an aliphatic amino acid residue

<220>

<223> At position 6, Xaa is an aliphatic amino acid residue

<220>

<223> At position 8, Xaa is a basic amino acid residue

<400> 318

```
Arg Pro Xaa Xaa Pro Xaa Arg Xaa Ser Xaa Pro
                 5
 1
<210> 319
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<400> 319
Pro Pro Val Pro Pro Arg Pro Xaa Xaa Thr Leu
                5
<210> 320
<211> 7
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: SH3 ANTAGONIST
<223> At positions 1, 3 and 6, Xaa is an aliphatic
      amino acid residue
<400> 320
Xaa Pro Xaa Leu Pro Xaa Lys
- 1
<210> 321
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SH3 ANTAGONIST
```

<223> At position 1, Xaa is a basic amino acid residue

<220> ...

```
<220>
<223> At position 2, Xaa is an aromatic amino acid
      residue
<400> 321
Xaa Xaa Asp Xaa Pro Leu Pro Xaa Leu Pro
                  5
<210> 322
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: INHIBITOR OF
      PLATELET AGGREGATION
<400> 322
Cys Xaa Xaa Arg Gly Asp Cys
 1
<210> 323
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SRC ANTAGONIST
<400> 323
Arg Pro Leu Pro Pro Leu Pro
                  5
<210> 324
<211> 6
<212> PRT
<213> Artificial Sequence
```

<223> Description of Artificial Sequence: SRC ANTAGONIST -

<400> 324

```
Pro Pro Val Pro Pro Arg
<210> 325
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:ANTI-CANCER
      PEPTIDE
<400> 325
Xaa Phe Xaa Asp Xaa Trp Xaa Xaa Leu Xaa Xaa
1 5
<210> 326
<211> 20
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:p16-MIMETIC
      PEPTIDE
<400> 326
Lys Ala Cys Arg Arg Leu Phe Gly Pro Val Asp Ser Glu Gln Leu Ser
                                                      15
                                10
                  5
 1
Arg Asp Cys Asp
             20
 <210> 327
 <211> 20
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:p16-MIMETIC
      PEPTIDE
```

136

<400> 327

```
Arg Glu Arg Trp Asn Phe Asp Phe Val Thr Glu Thr Pro Leu Glu Gly
                 5
                                    10
Asp Phe Ala Trp
             20
<210> 328
<211> 20
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:p16-MIMETIC
      PEPTIDE
<400> 328
Lys Arg Arg Gln Thr Ser Met Thr Asp Phe Tyr His Ser Lys Arg Arg
                                    10
                 5
Leu Ile Phe Ser
           20
<210> 329
<211> 20
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: SH3 ANTAGONIST
Thr Ser Met Thr Asp Phe Tyr His Ser Lys Arg Arg Leu Ile Phe Ser
                  5
                                    10
Lys Arg Lys Pro
             20
```

<211> 5 <212> PRT ... <213> Artificial Sequence

<210> 330

PCT/US99/25044

WO 00/24782 <220> <223> Description of Artificial Sequence:p16-MIMETIC PEPTIDE <400> 330 Arg Arg Leu Ile Phe 1 <210> 331 <211> 36 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence:p16-MIMETIC PEPTIDE <400> 331 Lys Arg Arg Gln Thr Ser Ala Thr Asp Phe Tyr His Ser Lys Arg Arg 10 5 1 Leu Ile Phe Ser Arg Gln Ile Lys Ile Trp Phe Gln Asn Arg Arg Met 25 Lys Trp Lys Lys

35

<210> 332 <211> 24 <212> PRT <213> Artificial Sequence

<223> Description of Artificial Sequence:p16-MIMETIC PEPTIDE

<400> 332 Lys Arg Arg Leu Ile Phe Ser Lys Arg Gln Ile Lys Ile Trp Phe Gln 10

Asn Arg Arg Met Lys Trp Lys Lys ... 20

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<210> 333
<211> 8
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: POLYGLYCINE
     LINKER
<400> 333
Gly Gly Gly Lys Gly Gly Gly
          5
<210> 334
<211> 8
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: POLYGLYCINE
    LINKER
<400> 334
Gly Gly Gly Asn Gly Ser Gly Gly
         5
<210> 335
<211> 8
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: POLYGLYCINE
     LINKER
<400> 335
Gly Gly Gly Cys Gly Gly Gly
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<210> 336 <211> 5

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:FC PCR PRIMER

<400> 336

Gly Pro Asn Gly Gly 1 5

<210> 337

<211> 42

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC

<400> 337

Phe Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg. Gln Trp Leu

1 5 10 15

Ala Ala Arg Ala Gly Gly Gly Gly Gly Gly Gly Ile Glu Gly Pro 20 25 30

Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala
35 40

<210> 338

<211> 42

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC

<400> 338

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly
1 5 10 15

Gly Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu ... 20 .25 30 ....

Ala Ala Arg Ala Gly Gly Gly Gly Phe

35 40

<210> 339

<211> 50

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC

<400> 339

Phe Gly Gly Gly Gly Gly Gly Thr Tyr Ser Cys His Phe Gly Pro 1 5 10 15

Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys Pro Gln 35 40 45

Gly Gly

50

<210> 340

<211> 50

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EPO-MIMETIC

<400> 340

Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys
1 5 10 15

Pro Gln Gly Gly Gly Gly Gly Gly Gly Thr Tyr Ser Cys His Phe 20 25 · 30

Gly Pro Leu Thr Trp Val Cys Lys Pro Gln Gly Gly Gly Gly Gly Gly 35 40 45

Gly Phe ...

<210> 341

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<211> 28
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TPO-MIMETIC
     PEPTIDES
<400> 341
Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Ile Glu
                                   10
                5
Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala
<210> 342
<211> 29
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TPO-MIMETIC
<400> 342
Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Ile
1 5
                                  10
Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala
                              25
            20
<210> 343
<211> 30
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: TPO-MIMETIC
<400> 343 ...
Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly
                                  10
                5
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Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala
20 25 30

<210> 344

<211> 31

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC

<400> 344

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly

1 5 10 15

Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala 20 25 30

<210> 345

<211> 32

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC

<400> 345

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly
1 5 10 15

Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala 20 25 30

<210> 346

<211> 33

<212> PRT ...

<213> Artificial Sequence

<220> <223> Description of Artificial Sequence: TPO-MIMETIC <400> 346 Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg 25 Ala <210> 347 <211> 34 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: TPO-MIMETIC <400> 347 Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly 5 Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala 25 20 Arg Ala <210> 348 <211> 35 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: TPO-MIMETIC <400> 348 Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly **1**5 _ . 10 ··· 5 Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala

20 25 30

Ala Arg Ala 35

<210> 349

<211> 36

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC

<400> 349

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly
1 5 10 15

Gly Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu 20 25 30

Ala Ala Arg Ala 35

<210> 350

<211> 37

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDES

<400> 350

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly
1 5 10 15

Gly Gly Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp
20 25 30

Leu Ala Ala Arg Ala

35

<210> 351

<211> 38

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDES

<400> 351

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly
1 5 10 15

Gly Gly Gly Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln 20 25 30

Trp Leu Ala Ala Arg Ala 35

<210> 352

<211> 42

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDES

<400> 352

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly
1 5 10 15

Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala 35 40

<210> 353

<211> 32

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDES

<400> 353

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Pro 1 5 10 15

Asn Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala
20 25 30

<210> 354

<211> 36

<212> PRT

<213> Artificial Sequence

<220>

<400> 354

Ile Glu Gly Pro Thr Leu Arg Gln Cys Leu Ala Ala Arg Ala Gly Gly
1 5 10 15

Gly Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Cys Leu 20  $\cdot$  25 30

Ala Ala Arg Ala 35

<210> 355

<211> 36

<212> PRT

<213> Artificial Sequence

<220>

<400> 355 ...

Ile Glu Gly Pro Thr Leu Arg Gln Cys Leu Ala Ala Arg Ala Gly Gly

1 5 10 15

Gly Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Cys Leu 20 25 30

Ala Ala Arg Ala 35

<210> 356

<211> 36

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDES

<400> 356

Ile Glu Gly Pro Thr Leu Arg Gln Ala Leu Ala Ala Arg Ala Gly Gly
1 5 10 15

Gly Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Ala Leu 20 25 30

Ala Ala Arg Ala 35

<210> 357

<211> 36

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDES

<400> 357

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly

1 5 10 15

Gly Lys Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu 20 25 30

Ala Ala Arg Ala

35

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<210> 358
<211> 40
<212> PRT
<213> Artificial Sequence
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<223> Description of Artificial Sequence: TPO-MIMETIC
      PEPTIDES
<400> 358
Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly
                                     10
Gly Lys Asx Arg Ala Cys Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu
                                 25
Arg Gln Trp Leu Ala Ala Arg Ala
         35
<210> 359
<211> 36
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: TPO-MIMETIC
      PEPTIDES
<400> 359
Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly
                 5
Gly Cys Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu
             20
                                25
Ala Ala Arg Ala
         35
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<213> Artificial Sequence

<220>

<400> 360

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly

1 5 10 15 .

Gly Lys Pro Glu Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg
20 25 30

Gln Trp Leu Ala Ala Arg Ala 35

<210> 361

<211> 39

<212> PRT

<213> Artificial Sequence

<220>

<400> 361

Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly

1 5 10 15

Gly Cys Pro Glu Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg
20 25 30

Gin Trp Leu Ala Ala Arg Ala 35

<210> 362

<211> 36

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDES

<400> 362 Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly 5 Gly Asn Gly Ser Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu . 25 Ala Ala Arg Ala 35 <210> 363 . <211> 36 <212> PRT <213> Artificial Sequence <223> Description of Artificial Sequence: TPO-MIMETIC PEPTIDES <400> 363 Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly 10 Gly Cys Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu 25 Ala Ala Arg Ala 35 <210> 364 <211> 57 <212> DNA <213> Artificial Sequence <220>

<223> Description of Artificial Sequence:Fc-TMP PCR PRIMER

<400> 364 aaaaaaggat cctcgagatt aagcacgagc agccagccac tgacgcagag tcggacc

<210> 365 <211> 39

<212>	DNA		
<213>	Artificial Sequence		
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<220>			
<223>	Description of Artificial Sequence:Fc-TMP PCR		
	PRIMER		
<400>	365		
	ggag gtggtggtat cgaaggteeg actetgegt		39
<210>	366		
<211>			
<212>			
	Artificial Sequence		
-013-	*** ***********************************		
<220>			
	Description of Artificial Sequence: INTEGRIN		
~223~	BINDING PEPTIDE		
	PINDING PERILUE		
<400>	366		
			42
cagtg	getgg etgetegtge ttaatetega ggateetttt tt		-
101.05	200		
<210>			
<211>			
<212>			
<213>	Artificial Sequence		
<220>			
<223>	Description of Artificial Sequence:Fc-TMP		•
<400>	367		60
	tggag gtggtggtat cgaaggtccg actctgcgtc agtggctggc	Egetegteet	81
taatc	tcgag gatccttttt t		91
	•		
<210>	368		
<211>	52		
<212>	DNA		
<213>	Artificial Sequence		
<220>			
<223>	Description of Artificial Sequence: Fc-TMP		
<400>			
ttcga	tacca ccacctccac ctttacccgg agacagggag aggctcttct	<b>gc</b>	52

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<210> 369
<211> 60
<212> DNA
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<223> Description of Artificial Sequence:Fc-TMP-TMP
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aaaggtggag gtggtggtat cgaaggtccg actctgcgtc agtggctggc tgctcgtgct 60
<210> 370
<211> 48
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:FC PCR PRIMER
<400> 370
acctccacca ccagcacgag cagccagcca ctgacgcaga gtcggacc
                                                                  48
<210> 371
<211> 66
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:Fc-TMP-TMP
      OLIGONUCLEOTIDE
<400> 371
ggtggtggag gtggcggcgg aggtattgag ggcccaaccc ttcgccaatg gcttgcagca 60
                                                                  66
cgcgca
<210> 372
<211> 76
<212> DNA
<213> Artificial Sequence
<223> Description of Artificial Sequence:Fc-TMP-TMP
      OLIGONUCLEOTIDE
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<400> 372 aaaaaaagga tootogagat tatgogogtg otgoaagcoa ttggogaagg gttgggooot 60 caatacctcc gccgcc <210> 373 <211> 126 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:Fc-TNF ALPHA PCR PRIMER <220> <221> CDS <222> (1)..(126) <400> 373 aaa ggt gga ggt ggt atc gaa ggt ccg act ctg cgt cag tgg ctg Lys Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu 10 1 5 gct gct cgt gct ggt ggt ggt ggc ggc gga ggt att gag ggc cca 96 Ala Ala Arg Ala Gly Gly Gly Gly Gly Gly Gly Ile Glu Gly Pro 20 126 acc ctt cgc caa tgg ctt gca gca cgc gca Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala 40 35 <210> 374 <211> 42 <212> PRT <213> Artificial Sequence <223> Description of Artificial Sequence:Fc-TNF ALPHA PCR PRIMER Lys Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu 10 5 1 Ala Ala Arg Ala Gly Gly Gly Gly Gly Gly Gly Ile Glu Gly Pro 25

40

Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala

35

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<210> 375
<211> 39
<212> DNA
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<223> Description of Artificial Sequence:Fc-MMP
      INHIBITOR
<220>
<221> CDS
<222> (4)..(732)
<400> 375
                                                                  39
ttt ttt cat atg atc gaa ggt ccg act ctg cgt cag tgg
    Phe His Met Ile Glu Gly Pro Thr Leu Arg Gln Trp
                      5
<210> 376
<211> 12
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:Fc-MMP
      INHIBITOR
<400> 376
Phe His Met Ile Glu Gly Pro Thr Leu Arg Gln Trp
                  5
<210> 377
<211> 48.
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: MMP INHIBITOR
<220>
<221> CDS ...
<222> (4)..(753)
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<400> 377
age acg age age cag cea etg acg cag agt egg ace tte gat cat atg
   Thr Ser Ser Gln Pro Leu Thr Gln Ser Arg Thr Phe Asp His Met
                                     10
                   5
                          .
<210> 378
<211> 15
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:MMP INHIBITOR
     FC
<400> 378 ·
Thr Ser Ser Gln Pro Leu Thr Gln Ser Arg Thr Phe Asp His Met
        5 .
                                   10
<210> 379
<211> 45
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:TMP-TMP-Fc
     OLIGONUCLEOTIDE
<400> 379
                                                               45
ctggctgctc gtgctggtgg aggcggtggg gacaaaactc acaca
<210> 380
<211> 51
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: INTEGRIN
      BINDING PEPTIDE
<400> 380
ctggctgctc gtgctggcgg tggtggcgga gggggtggca ttgagggccc a
                                                          51
<210> 381 ...
<211> 54
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<212> DNA

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<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: INTEGRIN
      BINDING PEPTIDE
<400> 381
                                                                  54
aagccattgg cgaagggttg ggccctcaat gccacccct ccgccaccac cgcc
<210> 382
<211> 54
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:INTEGRIN
      BINDING PEPTIDE
<400> 382
                                                                  54
accettegee aatggettge ageaegegea gggggaggeg gtggggaeaa aact
<210> 383
<211> 27
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: INTEGRIN
      BINDING PEPTIDE
<400> 383
                                                                  27
cccaccgcct ccccctgcgc gtgctgc
<210> 384
<211> 189
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: INTEGRIN
      BINDING PEPTIDE
<220>
<221> CDS
<222> (10)..(189)
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<400> 384

ttttttcat atg atc gaa ggt ccg act ctg cgt cag tgg ctg gct gct cgt 51

Met Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg

1 5 10

gct ggc ggt ggc gga ggg ggt ggc att gag ggc cca acc ctt cgc 99
Ala Gly Gly Gly Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg
15 20 25 30

caa tgg ctg gct cgt gct ggt gga ggc ggt ggg gac aaa act ctg 147 Gln Trp Leu Ala Ala Arg Ala Gly Gly Gly Gly Asp Lys Thr Leu 35 40 45

gct gct cgt gct ggt gga ggc ggt ggg gac aaa act cac aca 189
Ala Ala Arg Ala Gly Gly Gly Gly Asp Lys Thr His Thr
50 55 60

<210> 385

<211> 60

<212> PRT

<213> Artificial Sequence

<223> Description of Artificial Sequence: INTEGRIN BINDING PEPTIDE

<400> 385

Met Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly

1 5 10 15

Gly Gly Gly Gly Gly Gly Ile Glu Gly Pro Thr Leu Arg Gln Trp
20 25 30

Leu Ala Arg Ala Gly Gly Gly Gly Gly Asp Lys Thr Leu Ala Ala 35 40 45

Arg Ala Gly Gly Gly Gly Asp Lys Thr His Thr 50 55 60

<210> 386

<211> 141

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: INTEGRIN

## BINDING PEPTIDE

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<400> 386
ctaattccgc tctcacctac caaacaatgc cccctgcaa aaaataaatt catataaaaa 60
acatacagat aaccatctgc ggtgataaat tatctctggc ggtgttgaca taaataccac 120
tggcggtgat actgagcaca t
<210> 387
<211> 55
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: INTEGRIN
      BINDING PEPTIDE
<400> 387
cgatttgatt ctagaaggag gaataacata tggttaacgc gttggaattc ggtac
                                                                  55
<210> 388
<211> 872
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: INTEGRIN
      BINDING PEPTIDE
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gttagatatt tatcccttgc ggtgatagat tgagcacatc gatttgattc tagaaggagg 120
gataatatat gagcacaaaa aagaaaccat taacacaaga gcagcttgag gacgcacgtc 180
geettaaage aatttatgaa aaaaagaaaa atgaaettgg ettateeeag gaatetgteg 240
cagacaagat ggggatgggg cagtcaggcg ttggtgcttt atttaatggc atcaatgcat 300
taaatgctta taacgccgca ttgcttacaa aaattctcaa agttagcgtt gaagaattta 360
gcccttcaat cgccagagaa tctacgagat gtatgaagcg gttagtatgc agccgtcact 420
tagaagtgag tatgagtacc ctgttttttc tcatgttcag gcagggatgt tctcacctaa 480
gcttagaacc tttaccaaag gtgatgcgga gagatgggta agcacaacca aaaaagccag 540
tgattctgca ttctggcttg aggttgaagg taattccatg accgcaccaa caggctccaa 600
gccaagcttt cctgacggaa tgttaattct cgttgaccct gagcaggctg ttgagccagg 660
tgatttctgc atagccagac ttgggggtga tgagtttacc ttcaagaaac tgatcaggga 720
tageggteag gtgtttttae aaccactaaa eccacagtae ecaatgatee catgeaatga 780
gagttgttcc gttgtgggga aagttatcgc tagtcagtgg cctgaagaga cgtttggctg 840
                                                                   872
 atagactagt ggatccacta gtgtttctgc cc
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<210> 389
<211> 1197
<212> DNA
<213> Artificial Sequence
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<223> Description of Artificial Sequence: INTEGRIN
      BINDING PEPTIDE
<400> 389
ggcggaaacc gacgtccatc gaatggtgca aaacctttcg cggtatggca tgatagcgcc 60
cggaagagag tcaattcagg gtggtgaatg tgaaaccagt aacgttatac gatgtcgcag 120
agtatgccgg tgtctcttat cagaccgttt cccgcgtggt gaaccaggcc agccacgttt 180
ctgcgaaaac gcgggaaaaa gtcgaagcgg cgatggcgga gctgaattac attcccaacc 240
gcgtggcaca acaactggcg ggcaaacagt cgctcctgat tggcgttgcc acctccagtc 300
tggccctgca cgcgccgtcg caaattgtcg cggcgattaa atctcgcgcc gatcaactgg 360
gtgccagcgt ggtggtgtcg atggtagaac gaagcggcgt cgaagcctgt aaagcggcgg 420
tgcacaatct tctcgcgcaa cgcgtcagtg ggctgatcat taactatccg ctggatgacc 480
aggatgccat tgctgtggaa gctgcctgca ctaatgttcc ggcgttattt cttgatgtct 540
ctgaccagac acccatcaac agtattattt tctcccatga agacggtacg cgactgggcg 600
tggagcatct ggtcgcattg ggtcaccagc aaatcgcgct gttagcgggc ccattaagtt 660
ctgtctcggc gcgtctgcgt ctggctggct ggcataaata tctcactcgc aatcaaattc 720
agecgatage ggaacgggaa ggcgaetgga gtgccatgte eggtttteaa caaaccatge 780
aaatgctgaa tgagggcatc gttcccactg cgatgctggt tgccaacgat cagatggcgc. 840
tgggcgcaat gcgcgccatt accgagtccg ggctgcgcgt tggtgcggat atctcggtag 900
tgggatacga cgataccgaa gacagctcat gttatatccc gccgttaacc accatcaaac 960
aggattttcg cctgctgggg caaaccagcg tggaccgctt gctgcaactc tctcagggcc 1020
aggeggtgaa gggcaateag etgttgeeeg teteaetggt gaaaagaaaa accaeeetgg 1080
cgcccaatac gcaaaccgcc tctccccgcg cgttggccga ttcattaatg cagctggcac 1140
gacaggtttc ccgactggaa agcggacagt aaggtaccat aggatccagg cacagga
<210> 390
<211> 61
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:Fc-EMP
      OLIGONUCLEOTIDE
<400> 390
tatgaaaggt ggaggtggtg gtggaggtac ttactcttgc cacttcggcc cgctgacttg 60
g
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<210> 391 <211> 72

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<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:Fc-EMP
      OLIGONUCLEOTIDE
<400> 391
cggtttgcaa acccaagtca gcgggccgaa gtggcaagag taagtacctc caccaccacc 60
 tccacctttc at
 <210> 392
 <211> 57
 <212> DNA
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:Fc-EMP
       OLIGONUCLEOTIDE
 <400> 392
 gtttgcaaac cgcagggtgg cggcggcggc ggcggtggta cctattcctg tcatttt
 <210> 393
 <211> 60
 <212> DNA
 <213> Artificial Sequence
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       OLIGONUCLEOTIDE
 <400> 393
 ccaggtcagc gggccaaaat gacaggaata ggtaccaccg ccgccgccgc cgccaccctg 60
 <210> 394
 <211> 118
 <212> DNA
 <213> Artificial Sequence
<220>
  <223> Description of Artificial Sequence:Fc-EMP PCR
        TEMPLATE
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<220>

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<221> CDS <222> (2)..(118)

<400> 394

t atg aaa ggt gga ggt ggt ggt gga ggt act tac tct tgc cac ttc ggc 49 Met Lys Gly Gly Gly Gly Gly Gly Thr Tyr Ser Cys His Phe Gly

ccg ctg act tgg gtt tgc aaa ccg cag ggt ggc ggc ggc ggc ggt Pro Leu Thr Trp Val Cys Lys Pro Gln Gly Gly Gly Gly Gly Gly 20

ggt acc tat tcc tgt cat ttt Gly Thr Tyr Ser Cys His Phe 35

118

<210> 395

<211> 39

<212> PRT

<213> Artificial Sequence

<223> Description of Artificial Sequence:Fc-EMP PCR TEMPLATE

<400> 395

Met Lys Gly Gly Gly Gly Gly Gly Thr Tyr Ser Cys His Phe Gly 10 5

Pro Leu Thr Trp Val Cys Lys Pro Gln Gly Gly Gly Gly Gly Gly 25 20

Gly Thr Tyr Ser Cys His Phe 35

<210> 396

<211> 61

<212> DNA

<213> Artificial Sequence

<223> Description of Artificial Sequence:Fc-EMP PCR PRIMER

<400> 396

gcagaagagc ctctccctgt ctccgggtaa aggtggaggt ggtggtggag gtacttactc 60 61

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<210> 397
<211> 40
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:Fc-EMP PCR
       PRIMER
<400> 397
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ctaattggat ccacgagatt aaccaccctg cggtttgcaa
<210> 398
 <211> 22
 <212> DNA
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:Fc PRIMER
 <400> 398
                                                                   22
 aacataagta cctgtaggat cg
 <210> 399
 <211> 61
 <212> DNA
<213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:Fc PRIMER
 <400> 399
 agagtaagta cctccacca cacctccacc tttacccgga gacagggaga ggctcttctg 60
 <210> 400
 <211> 61
 <212> DNA
 <213> Artificial Sequence
 <220>
  <223> Description of Artificial Sequence: EMP-Fc
        OLIGONUCLEOTIDE
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<400> 400
ggcccgctga cctgggtatg taagccacaa gggggtgggg gaggcggggg gtaatctcga 60
<210> 401
<211> 50
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EMP-Fc
     OLIGONUCLEOTIDE
<400> 401
                                                              50
gatectegag attacecece geetececea ecceettgtg gettacatae
<210> 402
<211> 118
<212> DNA
<213> Artificial Sequence
<223> Description of Artificial Sequence: EMP-Fc PCR
      TEMPLATE
<220>
<221> CDS
<222> (1)..(108)
<400> 402
gtt tgc aaa ccg cag ggt ggc ggc ggc ggc ggt ggt acc tat tcc
                                                                 48
Val Cys Lys Pro Gln Gly Gly Gly Gly Gly Gly Gly Thr Tyr Ser
                                                       15
tgt cat ttt ggc ccg ctg acc tgg gta tgt aag cca caa ggg ggt ggg
                                                                 96
Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys Pro Gln Gly Gly
                                25
             20
                                                                 118
gga ggc ggg ggg taatctcgag
Gly Gly Gly Gly
        35
<210> 403
<211> 36
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<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: EMP-Fc PCR
<400> 403
Val Cys Lys Pro Gln Gly Gly Gly Gly Gly Gly Gly Thr Tyr Ser
                                    10
Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys Pro Gln Gly Gly
                        . 25
             20
Gly Gly Gly Gly
        35
<210> 404
<211> 39
<212> DNA
<213> Artificial Sequence
<223> Description of Artificial Sequence: EMP-Fc PCR
      PRIMER
<400> 404
                                                                39
ttatttcata tgaaaggtgg taactattcc tgtcatttt
<210> 405
<211> 43
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EMP-Fc PCR
      PRIMER
<400> 405
                                                                43
tggacatgtg tgagttttgt ccccccgcc tcccccaccc cct
<210> 406
<211> 43
<212> DNA
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<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:Fc PRIMER

<400> 406

agggggtggg ggaggcgggg gggacaaaac tcacacatgt cca

43

<210> 407

<211> 20

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:Fc PRIMER

<400> 407

gttattgctc agcggtggca

20

<210> 408

<211> 60

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:EMP-EMP-Fc
 OLIGONUCLEOTIDE

<400> 408

ttttttatcg atttgattct agatttgagt tttaactttt agaaggagga ataaaatatg 60

<210> 409

<211> 41

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:EMP-EMP-Fc
OLIGONUCLEOTIDE

<400> 409

taaaagttaa aactcaaatc tagaatcaaa tcgataaaaa a

41

<210> 410 ***

<211> 51

<212> DNA

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<213> Artificial Sequence
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     OLIGONUCLEOTIDE
<400> 410
ggaggtactt actcttgcca cttcggcccg ctgacttggg tttgcaaacc g
                                                                 51
<210> 411
<211> 55
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EMP-EMP-Fc
     OLIGONUCLEOTIDE
<400> 411
agtcagcggg ccgaagtggc aagagtaagt acctcccata ttttattcct ccttc
                                                                 55
<210> 412
<211> 60
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EMP-EMP-Fc
      OLIGONUCLEOTIDE
<400> 412
cagggtggcg gcggcggcgg cggtggtacc tattcctgtc attttggccc gctgacctgg 60
<210> 413
<211> 60
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EMP-EMP-Fc
      OLIGONUCLEOTIDE
<400> 413
aaaatgacag gaataggtac caccgccgcc gccgccgcca ccctgcggtt tgcaaaccca 60
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<210> 414
<211> 57
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EMP-EMP-Fc
      OLIGONUCLEOTIDE
<400> 414
gtatgtaagc cacaaggggg tgggggaggc gggggggaca aaactcacac atgtcca
                                                                57
<210> 415
<211> 60
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EMP-EMP-Fc
      OLIGONUCLEOTIDE
<400> 415
agttttgtcc cccccgcctc ccccaccccc ttgtggctta catacccagg tcagcgggcc 60
<210> 416
<211> 228
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EMP-EMP-Fc PCR
      TEMPLATE
<220>
<221> CDS
<222> (58)..(228)
<400> 416
ttttttatcg atttgattct agatttgagt tttaactttt agaaggagga ataaaat
                                                                57
atg gga ggt act tac tct tgc cac ttc ggc ccg ctg act tgg gtt tgc
Met Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys
                                                    1-5
                          . 10
        ··· 5
aaa ccg cag ggt ggc ggc ggc ggc ggt ggt acc tat tcc tgt cat
                                                                153
```

Lys Pro Gln Gly Gly Gly Gly Gly Gly Gly Thr Tyr Ser Cys His
20 25 30

ttt ggc ccg ctg acc tgg gta tgt aag cca caa ggg ggt ggg gga ggc 201 Phe Gly Pro Leu Thr Trp Val Cys Lys Pro Gln Gly Gly Gly Gly Gly 35 40 45

ggg ggg gac aaa act cac aca tgt cca 228
Gly Gly Asp Lys Thr His Thr Cys Pro
50 55

<210> 417

<211> 57

<212> PRT

<213> Artificial Sequence

<223> Description of Artificial Sequence: EMP-EMP-Fc PCR TEMPLATE

<400> 417

Met Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys 1 5 10 15

Lys Pro Gln Gly Gly Gly Gly Gly Gly Gly Thr Tyr Ser Cys His 20 25 30

Phe Gly Pro Leu Thr Trp Val Cys Lys Pro Gln Gly Gly Gly Gly Gly 35 40 45

Gly Gly Asp Lys Thr His Thr Cys Pro 50 55

<210> 418

<211> 40

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:Fc-EMP-EMP PCR
PRIMER

<400> 418

ctaattggat cctcgagatt aaccccttg tggcttacat

40

<210> 419

<211> 72

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EPO-MIMETIC PEPTIDE

<400> 419

Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa 70

<210> 420

<211> 62

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EPO-MIMETIC PEPTIDE

<400> 420

Xaa Tyr Xaa Xaa Cys Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa Gly Pro

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<210> 421
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EPO-MIMETIC
      PEPTIDE
<220>
<223> At position 2, Xaa is R, H, L or W
<220>
<223> At position 3, Xaa is M, F or I
<220>
<223> At position 6, Xaa is any of the 20 genetically
      encoded amino acid residues or a D-stereoisomer
      thereof
<220>
<223> At position 9, Xaa is D, E, I, L or V
<400> 421
Cys Xaa Xaa Gly Pro Xaa Thr Trp Xaa Cys
                 5
<210> 422
<211> 19
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EPO-MIMETIC
      PEPTIDE
<400> 422
Gly Gly Thr Tyr Ser Cys His Gly Pro Leu Thr Trp Val Cys Lys Pro
Gln Gly Gly
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<210> 423

```
<211> 19
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EPO-MIMETIC
     PEPTIDE
<400> 423
Val Gly Asn Tyr Met Ala His Met Gly Pro Ile Thr Trp Val Cys Arg
                                     10
Pro Gly Gly
<210> 424
<211> 18
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EPO-MIMETIC
      PEPTIDE
<400> 424
Gly Gly Pro His His Val Tyr Ala Cys Arg Met Gly Pro Leu Thr Trp
                                     10
Ile Cys
<210> 425
<211> 18
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: EPO-MIMETIC
       PEPTIDE
<400> 425
Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys
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1 5 10 15

Pro Gln

<210> 426

<211> 20

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EPO-MIMETIC PEPTIDE

<400> 426

Gly Gly Leu Tyr Ala Cys His Met Gly Pro Met Thr Trp Val Cys Gln
1 5 10 15

Pro Leu Arg Gly 20

<210> 427

<211> 22

<212> PRT

<213> Artificial Sequence

<220>

<400> 427

Thr Ile Ala Gln Tyr Ile Cys Tyr Met Gly Pro Glu Thr Trp Glu Cys
1 5 10 15

Arg Pro Ser Pro Lys Ala 20

<210> 428

<211> 13

<212> PRT...

<213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: EPO-MIMETIC
      PEPTIDE
<400> 428
Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys
                                    10
                5
<210> 429
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EPO MIMETIC
      PEPTIDE
<400> 429
 Tyr Cys His Phe Gly Pro Leu Thr Trp Val Cys
                                      10
                  5
 <210> 430
 <211> 17
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:UKR ANTAGONIST
       PEPTIDE
 <400> 430
 Ala Glu Pro Val Tyr Gln Tyr Glu Leu Asp Ser Tyr Leu Arg Ser Tyr
                   5
                                     10
 Tyr
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<210> 431 <211> 17

<212> PRT

<213> Artificial Sequence

<220> <223> Description of Artificial Sequence: UKR ANTAGONIST PEPTIDE <400> 431 Ala Glu Leu Asp Leu Ser Thr Phe Tyr Asp Ile Gln Tyr Leu Leu Arg 10 Thr <210> 432 <211> 17 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence:UKR ANTAGONIST PEPTIDE <400> 432 Ala Glu Phe Phe Lys Leu Gly Pro Asn Gly Tyr Val Tyr Leu His Ser 1 . 5 10 Ala <210> 433 <211> 11 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence:UKR ANTAGONIST PEPTIDE

<210> 434 <211> 17

Phe Lys Leu Xaa Xaa Kaa Gly Tyr Val Tyr Leu

5

```
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: UKR ANTAGONIST
      PEPTIDE
<400> 434
Ala Glu Ser Thr Tyr His His Leu Ser Leu Gly Tyr Met Tyr Thr Leu
                 5
                                    10
Asn
<210> 435
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: UKR ANTAGONIST
      PEPTIDE
<400> 435
Tyr His Xaa Leu Xaa Xaa Gly Tyr Met Tyr Thr
                  5
<210> 436
<211> 6
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: MCA/MCP
      INHIBITOR
<400> 436
Arg Asn Arg Gln Lys Thr
                 5
 1
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<210> 437 <211> 4

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<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:MCA/MCP
      INHIBITOR
<400> 437
Arg Asn Arg Gln
 1
<210> 438
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:MCA/MCP
      INHIBITOR
<400> 438
Arg Asn Arg Gln Lys
 1
<210> 439
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: MCA/MCP
      INHIBITOR
<400> 439
Asn Arg Gln Lys Thr
  1
<210> 440
<211> 4
<212> PRT "
<213> Artificial Sequence
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<220>
<223> Description of Artificial Sequence:MCA/MCP
      INHIBITOR
<400> 440
Arg Gln Lys Thr
 1
<210> 441
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial
      Sequence: INTEGRIN-BINDING PEPTIDE
<400> 441
Arg Xaa Glu Thr Xaa Trp Xaa
  1
<210> 442
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial
      Sequence: INTEGRIN-BINDING PEPTIDE
<400> 442
Arg Xaa Glu Thr Xaa Trp Xaa
                 5
<210> 443
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial
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Sequence: INTEGRIN-BINDING PEPTIDE

```
<400> 443
Arg Gly Asp Gly Xaa
1 5
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<210> 444

<211> 7

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial
 Sequence:INTEGRIN-BINDING PEPTIDE

<400> 444

Cys Arg Gly Asp Gly Xaa Cys 1 5

<210> 445

<211> 9

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial
 Sequence:INTEGRIN-BINDING PEPTIDE

<400> 445

Cys Xaa Xaa Arg Leu Asp Xaa Xaa Cys 1 5

<210> 446

<211> 9

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial
 Sequence:INTEGRIN-BINDING PEPTIDE

<400> 446

Cys Ala Arg Arg Leu Asp Ala Pro Cys

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1 5

<210> 447

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<211> 9

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial
 Sequence:INTEGRIN-BINDING PEPTIDE

<400> 447

Cys Pro Ser Arg Leu Asp Ser Pro Cys

<210> 448

<211> 9

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:INTEGRIN-BINDING PEPTIDE

<400> 448

Xaa Xaa Xaa Arg Gly Asp Xaa Xaa Xaa 1 5

<210> 449

<211> 9

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:INTEGRIN-BINDING PEPTIDE

<400> 449

Cys Xaa Cys Arg Gly Asp Cys Xaa Cys

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<210> 450
<211> 9
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial
      Sequence: INTEGRIN-BINDING PEPTIDE
<400> 450
Cys Asp Cys Arg Gly Asp Cys Phe Cys
<210> 451
<211> 9
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial
      Sequence: INTEGRIN-BINDING PEPTIDE
<400> 451
Cys Asp Cys Arg Gly Asp Cys Leu Cys
          5
<210> 452
<211> 9
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial
      Sequence: INTEGRIN-BINDING PEPTIDE
<400> 452
Cys Leu Cys Arg Gly Asp Cys Ile Cys
  1
                  5
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<210> 453 <211> 8

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<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial
     Sequence: INTEGRIN-BINDING PEPTIDE
<400> 453
Xaa Xaa Asp Asp Xaa Xaa Xaa
                 5
<210> 454
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial
      Sequence: INTEGRIN-BINDING PEPTIDE
<400> 454
Xaa Xaa Xaa Asp Asp Xaa Xaa Xaa Xaa
                 5
<210> 455
<211> 8
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial
      Sequence: INTEGRIN-BINDING PEPTIDE
<400> 455
Cys Trp Asp Asp Gly Trp Leu Cys
                5
<210> 456
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<210> 456
<211> 9
<212> PRT --<213> Artificial Sequence

<220> <223> Description of Artificial Sequence: INTEGRIN-BINDING PEPTIDE <400> 456 Cys Trp Asp Asp Leu Trp Trp Leu Cys <210> 457 <211> 8 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: INTEGRIN-BINDING PEPTIDE <400> 457 Cys Trp Asp Asp Gly Leu Met Cys 5 <210> 458 <211> 8 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: INTEGRIN-BINDING PEPTIDE <400> 458 Cys Trp Asp Asp Gly Trp Met Cys 5 . <210> 459 <211> 9 <212> PRT <213> Artificial Sequence <220>

<223> Description of Artificial

Sequence: INTEGRIN-BINDING PEPTIDE

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<400> 459
Cys Ser Trp Asp Asp Gly Trp Leu Cys
1 5
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<210> 460

<211> 9

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial
 Sequence:INTEGRIN-BINDING PEPTIDE

<400> 460

Cys Pro Asp Asp Leu Trp Trp Leu Cys
1 5

<210> 461

<211> 40

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:EPO-MIMETIC
 PEPTIDE

<400> 461

Xaa Xaa Xaa Xaa Xaa Xaa Xaa 40

<210> 462

<211> 16

<212> PRT---

<213> Artificial Sequence

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10

15

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5

qaA

<210> 464 <211> 19 <212> PRT <213> Artificial Sequence <220>

<223> Description of Artificial Sequence: SELECTIN ANTAGONIST PEPTIDE

<400> 464 Arg Lys Asn Asn Lys Thr Trp Thr Trp Val Gly Thr Lys Lys Ala Leu 10

Thr Asn Glu

<210> 465 <211> 13

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SELECTIN ANTAGONIST PEPTIDE

<400> 465

Lys Lys Ala Leu Thr Asn Glu Ala Glu Asn Trp Ala Asp 1 5 10

<210> 466

<211> 16

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SELECTIN ANTAGONIST PEPTIDE

<400> 466

Cys Gln Xaa Arg Tyr Thr Asp Leu Val Ala Ile Gln Asn Lys Xaa Glu 1 5 10 15

<210> 467

<211> 19

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SELECTIN ANTAGONIST PEPTIDE

<400> 467

Arg Lys Xaa Asn Xaa Xaa Trp Thr Trp Val Gly Thr Xaa Lys Xaa Leu 1 5 10 15

Thr Glu Glu

<210> 468

<211> 17

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<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SELECTIN
      ANTAGONIST PEPTIDE
<400> 468
Ala Glu Asn Trp Ala Asp Gly Glu Pro Asn Asn Lys Xaa Asn Xaa Glu
                                    10
Asp
<210> 469
<211> 16
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SELECTIN
   ANTAGONIST PEPTIDE
<400> 469
Cys Xaa Xaa Xaa Tyr Thr Xaa Leu Val Ala Ile Gln Asn Lys Xaa Glu
                                    10
                 5
<210> 470
<211> 19
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SELECTIN
      ANTAGONIST PEPTIDE
 <400> 470
Arg Lys Xaa Xaa Xaa Trp Xaa Trp Val Gly Thr Xaa Lys Xaa Leu
                                   10
```

Thr Xaa Glu

1

5

```
<210> 471
<211> 16
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SELECTIN
      ANTAGONIST PEPTIDE
<400> 471
Ala Xaa Asn Trp Xaa Xaa Xaa Glu Pro Asn Asn Xaa Xaa Xaa Glu Asp
                                     10
<210> 472
<211> 13
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SELECTIN
      ANTAGONIST PEPTIDE
<400> 472
Xaa Lys Xaa Lys Thr Xaa Glu Ala Xaa Asn Trp Xaa Xaa
                5
<210> 473
<211> 12
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: SOMATOSTATIN/
      CORTISTATIN-MIMETIC PEPTIDE
<220>
<223> At position 1, Xaa is asp-arg-met-pro-cys,
      arg-met-pro-cys, met-pro-cys, pro-cys, or cys
<220>
<223> At position 2, Xaa is arg or lys
<220>
```

```
<223> At position 10, Xaa is ser or thr
 <220>
 <223> At position 12, xaa is cys:lys or cys
 <400> 473
 Xaa Xaa Asn Phe Phe Trp Lys Thr Phe Xaa Ser Xaa
                   5
 <210> 474
 <211> 18
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: SOMATOSTATIN/
       CORTISTATIN-MIMETIC PEPTIDE
 <400> 474
 Asp Arg Met Pro Cys Arg Asn Phe Phe Phe Trp Lys Thr Phe Ser Ser
                                      10
 Cys Lys
 <210> 475
<211> 15
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: SOMATOSTATIN/
       CORTISTATIN-MIMETIC PEPTIDE
 <400> 475
 Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys Lys
                   5
                                     .10
 <210> 476
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<211> 13 ... <212> PRT

<213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: SOMATOSTATIN/
      CORTISTATIN-MIMETIC PEPTIDE
<400> 476
Cys Arg Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys Lys
 1 . 5
<210> 477
<211> 16
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: SOMATOSTATIN/
      CORTISTATIN-MIMETIC PEPTIDE
<400> 477
Asp Arg Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys
                                    10
<210> 478
<211> 14
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SOMATOSTATIN/
      CORTISTATIN MIMETIC PEPTIDE
<400> 478
Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys
<210> 479
<211> 12
<212> PRT
<213> Artificial Sequence
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<223> Description of Artificial Sequence: SOMATOSTATIN/

<220>

## CORTISTATIN MIMETIC PEPTIDE

<400> 479
Cys Arg Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys
1 5 10

<210> 480

<211> 16

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SOMATOSTATIN/ CORTISTATIN MIMETIC PEPTIDE

<400> 480

Asp Arg Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys

1 5 10 15

<210> 481

<211> 15

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SOMATOSTATIN/ CORTISTATIN MIMETIC PEPTIDE

<400> 481

Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys Lys
1 5 10 15

<210> 482

<211> 13

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: SOMATOSTATIN/ CORTISTATIN MIMETIC PEPTIDE

<400> 482

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Cys Lys Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys Lys
                                    10
                5
<210> 483
<211> 16
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SOMATOSTATIN/
      CORTISTATIN MIMETIC PEPTIDE
<400> 483
Asp Arg Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys
                                     10
                  5
<210> 484
<211> 14
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:Fc-TMP
<400> 484
Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys
                                     10
                 5
 <210> 485
 <211> 12
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: SOMATOSTATIN/
       CORTISTATIN MIMETIC PEPTIDE
 Cys Lys Asn Phe Phe Trp Lys Thr Phe Ser Ser Cys
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1 ... 5

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<210> 486
<211> 17
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SOMATOSTATIN/
      CORTISTATIN MIMETIC PEPTIDE
<400> 486
Asp Arg Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys
                  5
                                    10
Lys
<210> 487
<211> 15
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SOMATOSTATIN/
     CORTISTATIN MIMETIC PEPTIDE
<400> 487
Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys Lys
                 5
                                     10
<210> 488
<211> 13
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: SOMATOSTATIN/
      CORTISTATIN MIMETIC PEPTIDE
<400> 488
Cys Arg Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys Lys
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5

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<210> 489
<211> 16
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SOMATOSTATIN/
     CORTISTATIN MIMETIC PEPTIDE
<400> 489
Asp Arg Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys
                                   10
                5
<210> 490
<211> 14
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SOMATOSTATIN/
      CORTISTATIN MIMETIC PEPTIDE
<400> 490
Met Pro Cys Arg Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys
                                    10
                 5
<210> 491
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SOMATOSTATIN/
      CORTISTATIN MIMETIC PEPTIDE
<400> 491
Cys Arg Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys
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<210> 492 <211> 17

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<212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: SOMATOSTATIN/
       CORTISTATIN MIMETIC PEPTIDE
 <400> 492
 Asp Arg Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys
                    5
                                     10
 Lys
 <210> 493
  <211> 15
 <212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence: SOMATOSTATIN/
        CORTISTATIN MIMETIC PEPTIDE
  <400> 493
  Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys Lys
                                      10
<210> 494
 <211> 13
  <212> PRT
  <213> Artificial Sequence
  <223> Description of Artificial Sequence: SOMATOSTATIN/
        CORTISTATIN MIMETIC PEPTIDE
  <400> 494
  Cys Lys Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys Lys
                   5
```

<210> 495 <211> 16

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<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SOMATOSTATIN/
      CORTISTATIN MIMETIC PEPTIDE
Asp Arg Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys
                5
                                    10
<210> 496
<211> 14
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SOMATOSTATIN/
    CORTISTATIN MIMETIC PEPTIDE
<400> 496
Met Pro Cys Lys Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys
                 5
<210> 497
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: SOMATOSTATIN/
      CORTISTATIN MIMETIC PEPTIDE
<400> 497
Cys Lys Asn Phe Phe Trp Lys Thr Phe Thr Ser Cys
<210> 498
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<211> 25 <212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:CAP37
MIMETIC/LPS BINDING PEPTIDE

<400> 498

Asn Gln Gly Arg His Phe Cys Gly Gly Ala Leu Ile His Ala Arg Phe 1 5 10 15

Val Met Thr Ala Ala Ser Cys Phe Gln
20 25

<210> 499

<211> 20

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: CAP37
MIMETIC/LPS BINDING PEPTIDE

<400> 499

Arg His Phe Cys Gly Gly Ala Leu Ile His Ala Arg Phe Val Met Thr
1 5 10 15

Ala Ala Ser Cys 20

<210> 500

<211> 27

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:CAP37
 MIMETIC/LPS BINDING PEPTIDE

<400> 500

Gly Thr Arg Cys Gln Val Ala Gly Trp Gly Ser Gln Arg Ser Gly Gly

1 5 10 15

Arg Leu Ser Arg Phe Pro Arg Phe Val Asn Val

**...** 20 . 25

```
<210> 501
<211> 18
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VEGF-ANTAGONIST
      PEPTIDE
<400> 501
Gly Glu Arg Trp Cys Phe Asp Gly Pro Arg Ala Trp Val Cys Gly Trp
                                     10
Glu Ile
<210> 502
<211> 18
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: VEGF ANTAGONIST
     PEPTIDE
Glu Glu Leu Trp Cys Phe Asp Gly Pro Arg Ala Trp Val Cys Gly Tyr
                  5
                                    10
                                                         15
Val Lys
<210> 503
<211> 33
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: ANTIPATHOGENIC
      PEPTIDE
<400> 503 .
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Gly Phe Phe Ala Leu Ile Pro Lys Ile Ile Ser Ser Pro Leu Phe Lys

1 5 10 15

Thr Leu Leu Ser Ala Val Gly Ser Ala Leu Ser Ser Gly Gly Gln 20 .25 30

Gln

<210> 504

<211> 33

<212> PRT

<213> Artificial Sequence

<220>

<220>

<223> At positions 7, 18 and 19, D amino acid residue

<400> 504

Gly Phe Phe Ala Leu Ile Pro Lys Ile Ile Ser Ser Pro Leu Phe Lys
1 5 10 15

Thr Leu Leu Ser Ala Val Gly Ser Ala Leu Ser Ser Ser Gly Gly Gln 20 25 30

Glu

<210> 505

<211> 22

<212> PRT

<213> Artificial Sequence

<220>

<220>

<223> At positions 18 and 19, D amino acid residues

<400> 505

Gly Phe Phe Ala Leu Ile Pro Lys Ile Ile Ser Ser Pro Leu Phe Lys

1 5 10 15

Thr Leu Leu Ser Ala Val 20

<210> 506

<211> 22

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<220>

<223> At positions 7, 18 and 19, D amino acid residues

<400> 506

Gly Phe Phe Ala Leu Ile Pro Lys Ile Ile Ser Ser Pro Leu Phe Lys
1 5 10 15

Thr Leu Leu Ser Ala Val

20

<210> 507

<211> 23

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<220>

<223> At positions 8, 19 and 20, D amino acid residues

<400> 507

Lys Gly Phe Phe Ala Leu Ile Pro Lys Ile Ile Ser Ser Pro Leu Phe
1 5 10 15

Lys Thr Leu Leu Ser Ala Val

20

```
<210> 508
<211> 24
 <212> PRT
<213> Artificial Sequence
<220>
 <223> Description of Artificial Sequence: VIP MIMETIC
       PEPTIDE
 <220>
 <223> At positions 9, 20 and 21, D amino acid residues
<400> 508
Lys Lys Gly Phe Phe Ala Leu Ile Pro Lys Ile Ile Ser Ser Pro Leu
                   5
                                      10
Phe Lys Thr Leu Leu Ser Ala Val
              20
<210> 509
<211> 24
<212> PRT
 <213> Artificial Sequence
<220>
 <223> Description of Artificial Sequence:VIP MIMETIC
       PEPTIDE
 <220>
 <223> At positions 9, 20 and 21, D amino acid residues
 <400> 509
 Lys Lys Gly Phe Phe Ala Leu Ile Pro Lys Ile Ile Ser Ser Pro Leu
                   5
                                      10
 Phe Lys Thr Leu Leu Ser Ala Val
              20
 <210> 510
 <211> 11
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<212> PRT-

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: VIP MIMETIC PEPTIDE

<220>

<223> At position 7, D amino acid residue

<400> 510

Gly Phe Phe Ala Leu Ile Pro Lys Ile Ile Ser 1 5 10

<210> 511

<211> 26

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 511

Gly Ile Gly Ala Val Leu Lys Val Leu Thr Thr Gly Leu Pro Ala Leu

1 5 10 15

Ile Ser Trp Ile Lys Arg Lys Arg Gln Gln 20 25

<210> 512

<211> 26

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<220>

<223> At positions 5, 8, 17 and 23, D amino acid residues

<400> 512

Gly Ile Gly Ala Val Leu Lys Val Leu Thr Thr Gly Leu Pro Ala Leu

1 5 10 15

Ile Ser Trp Ile Lys Arg Lys Arg Gln Gln 20 25

<210> 513

<211> 26

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<220>

<223> At positions 5, 8, 17 and 23, D amino acid residues

<400> 513

Gly Ile Gly Ala Val Leu Lys Val Leu Thr Thr Gly Leu Pro Ala Leu

1 5 10 15

Ile Ser Trp Ile Lys Arg Lys Arg Gln Gln 20 25

<210> 514

<211> 22

<212> PRT

<213> Artificial Sequence

<220>

<220>

<223> At positions 5, 8, 17 and 21, D amino acid residues

<400> 514

Gly Ile Gly Ala Val Leu Lys Val Leu Thr Thr Gly Leu Pro Ala Leu

1 5 10 15

Ile Ser Trp Ile Lys Arg

... 20

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<210> 515
<211> 19
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<220>
<223> At positions 2, 5, 14 and 18, D amino acid
      residues
<400> 515
Ala Val Leu Lys Val Leu Thr Thr Gly Leu Pro Ala Leu Ile Ser Trp
                                    10
                  5
Ile Lys Arg
<210> 516
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<220>
<223> At positions 3, 4, 8 and 10, D amino acid residues
.<400> 516
Lys Leu Leu Leu Leu Lys Leu Leu Leu Lys
                  5
  1
 <210> 517
 <211> 12
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:VIP MIMETIC
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PEPTIDE

<220>
<223> At positions 3, 4, 8 and 10, D amino acid residues
<400> 517
Lys Leu Leu Leu Lys Leu Leu Lys Leu Leu Lys
1 5 10

<210> 518 <211> 12 <212> PRT <213> Artificial Sequence

<220>
<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<220>
<223> At positions 3, 4, 8 and 10, D amino acid residues
<400> 518

Lys Leu Leu Lys Leu Lys Leu Lys Leu Lys

1 5 10

<210> 519
<211> 12
<212> PRT
<213> Artificial Sequence
<220>

<223> Description of Artificial Sequence: VIP MIMETIC PEPTIDE

<400> 519
Lys Lys Leu Leu Lys Leu Lys Leu Lys Leu Lys Lys
1 5 10

<210> 520 <211> 12 .... <212> PRT <213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: VIP MIMETIC
     PEPTIDE
<400> 520
Lys Leu Leu Lys Leu Leu Lys Leu Lys
                5
<210> 521
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
<400> 521
Lys Leu Leu Lys Leu Lys Leu Lys Leu Leu Lys
                 5
<210> 522
<211> 6
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
 <400> 522
 Lys Leu Leu Leu Lys
 <210> 523
 <211> 8
 <212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence: VIP MIMETIC
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PEPTIDE

<400> 523 Lys Leu Leu Leu Lys Leu Leu Lys 1 5

<210> 524 <211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 524

Lys Leu Leu Leu Lys Leu Lys Leu Lys Leu Leu Lys

1 5 10

<210> 525

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 525

Lys Leu Leu Leu Lys Leu Lys Leu Lys Leu Lys 1 5 10

<210> 526

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 526

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Lys Leu Leu Lys Leu Lys Leu Lys Leu Leu Lys
                5
- 1
<210> 527
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
<400> 527
Lys Ala Ala Ala Lys Ala Ala Lys Ala Ala Lys
                5
<210> 528
<211> 12
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 528
Lys Val Val Val Lys Val Val Lys Val Val Lys
                 5
<210> 529
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
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10

Lys Val Val Lys Val Lys Val Lys Val Val Lys

<400> 529 ...

. 5

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<210> 530
<211> 11
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 530
Lys Val Val Lys Val Lys Val Lys Val Lys
 1 . 5
<210> 531
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
Lys Val Val Lys Val Lys Val Lys Val Val Lys
                                   10
                 5
 1
 <210> 532
 <211> 6
 <212> PRT
<213> Artificial Sequence
 <223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
 <400> 532
 Lys Leu Ile Leu Lys Leu
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<210> 533

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<211> 6
<212> PRT
<213> Artificial Sequence
<220>
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      PEPTIDE
<400> 533
Lys Val Leu His Leu Leu
 1
<210> 534
<211> 6
<212> PRT
<213> Artificial Sequence
<220>
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      PEPTIDE
<400> 534
Leu Lys Leu Arg Leu Leu
 <210> 535
<211> 6
 <212> PRT
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   PEPTIDE
 <400> 535
 Lys Pro Leu His Leu Leu
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<210> 536 <211> 8 <212> PRT <213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 536
Lys Leu Ile Leu Lys Leu Val Arg
 1
<210> 537
<211> 8
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
       PEPTIDE
 <400> 537
 Lys Val Phe His Leu Leu His Leu
                   5
 <210> 538
 <211> 8
 <212> PRT
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  <220>
  <223> Description of Artificial Sequence: VIP MIMETIC
        PEPTIDE
  <400> 538
  His Lys Phe Arg Ile Leu Lys Leu
                    5
  <210> 539
  <211> 8
  <212> PRT
  <213> Artificial Sequence
   <220>
   <223> Description of Artificial Sequence: VIP MIMETIC
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PEPTIDE

<400> 539 Lys Pro Phe His Ile Leu His Leu 5

<210> 540

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 540

Lys Ile Ile Lys Ile Lys Ile Lys Ile Lys 10 5

<210> 541

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 541

Lys Ile Ile Ile Lys Ile Lys Ile Lys Ile Ile Lys 5

<210> 542

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: VIP MIMETIC PEPTIDE

<400> 542

Lys Ile Ile Ile Lys Ile Lys Ile Lys Ile Ile Lys
1 5 10

<210> 543

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC
PEPTIDE

<400> 543

Lys Ile Pro Ile Lys Ile Lys Ile Lys Ile Pro Lys
1 5 10

<210> 544

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 544

Lys Ile Pro Ile Lys Ile Lys Ile Lys Ile Val Lys
1 5 10

<210> 545

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<400> 545

Arg Ile Ile Ile Arg Ile Arg Ile Arg Ile Ile Arg

1 5 10

```
<210> 546
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
     PEPTIDE
<400> 546
Arg Ile Ile Arg Ile Arg Ile Arg Ile Ile Arg
 1 . 5
<210> 547
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
     PEPTIDE
<400> 547
Arg Ile Ile Ile Arg Ile Arg Ile Arg Ile Ile Arg
                                  10
                 5
<210> 548
 <211> 12
 <212> PRT
<213> Artificial Sequence
 <223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
 <400> 548
 Arg Ile Val Ile Arg Ile Arg Ile Arg Leu Ile Arg
                 5
  1
```

<210> 549

```
<211> 12
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: VIP MIMETIC
       PEPTIDE
 <400> 549
 Arg Ile Ile Val Arg Ile Arg Leu Arg Ile Ile Arg
   1
                   5
 <210> 550
 <211> 12
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:VIP MIMETIC
 PEPTIDE
 <400> 550
 Arg Ile Gly Ile Arg Leu Arg Val Arg Ile Ile Arg
  1
 <210> 551
<211> 12
 <212> PRT
 <213> Artificial Sequence
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 <223> Description of Artificial Sequence: VIP MIMETIC
       PEPTIDE
 <400> 551
 Lys Ile Val Ile Arg Ile Arg Ile Arg Leu Ile Arg
   1
                  5
```

<210> 552 <211> 12 ... <212> PRT

<213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 552
Arg Ile Ala Val Lys Trp Arg Leu Arg Phe Ile Lys
                 5
<210> 553
<211> 12
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 553
Lys Ile Gly Trp Lys Leu Arg Val Arg Ile Ile Arg
                 5
<210> 554
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 554
Lys Lys Ile Gly Trp Leu Ile Ile Arg Val Arg Arg
                 5
<210> 555
<211> 14
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: VIP MIMETIC
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PEPTIDE

<400> 555
Arg Ile Val Ile Arg Ile Arg Ile Arg Leu Ile Arg Ile Arg
1 5 10

<210> 556

<211> 14

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 556

Arg Ile Ile Val Arg Ile Arg Leu Arg Ile Ile Arg Val Arg
1 5 10

<210> 557

<211> 14

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 557

Arg Ile Gly Ile Arg Leu Arg Val Arg Ile Ile Arg Arg Val
1 5 10

<210> 558

<211> 16

<212> PRT

<213> Artificial Sequence

<220>

<400> 558

```
Lys Ile Val Ile Arg Ile Arg Ala Arg Leu Ile Arg Ile Arg Ile Arg
                   5
                                     10
                                                        15
  1
 <210> 559
 <211> 16
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:VIP MIMETIC
       PEPTIDE
 <400> 559
 Arg Ile Ile Val Lys Ile Arg Leu Arg Ile Ile Lys Lys Ile Arg Leu
                 5
                                    10
   1
 <210> 560
 <211> 16
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: VIP MIMETIC
       PEPTIDE
 Lys Ile Gly Ile Lys Ala Arg Val Arg Ile Ile Arg Val Lys Ile Ile
                  5
                                     10
  1
 <210> 561
 <211> 16.
<212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
 <400> 561
 Arg Ile Ile Val His Ile Arg Leu Arg Ile Ile His His Ile Arg Leu
```

5

10

```
<210> 562
 <211> 16
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:VIP MIMETIC
       PEPTIDE
 <400> 562
 His Ile Gly Ile Lys Ala His Val Arg Ile Ile Arg Val His Ile Ile
             5
 <210> 563
 <211> 16
 <212> PRT
 <213> Artificial Sequence
 <220>
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       PEPTIDE
 <400> 563
 Arg Ile Tyr Val Lys Ile His Leu Arg Tyr Ile Lys Lys Ile Arg Leu
  1 5
 <210> 564
 <211> 16
 <212> PRT
 '<213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:VIP MIMETIC
       PEPTIDE
 <400> 564
Lys Ile Gly His Lys Ala Arg Val His Ile Ile Arg Tyr Lys Ile Ile
                 5 10
  1
```

<210> 565

```
<211> 16
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 565
Arg Ile Tyr Val Lys Pro His Pro Arg Tyr Ile Lys Lys Ile Arg Leu
 1 5
                      10
<210> 566
<211> 16
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: VIP MIMETIC
    PEPTIDE
<400> 566
Lys Pro Gly His Lys Ala Arg Pro His Ile Ile Arg Tyr Lys Ile Ile
                5
<210> 567
 <211> 19
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
 <400> 567
 Lys Ile Val Ile Arg Ile Arg Ile Arg Leu Ile Arg Ile Arg
         5
Lys Ile Val
```

<210> 568

```
<211> 19
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
     PEPTIDE
<400> 568
Arg Ile Ile Val Lys Ile Arg Leu Arg Ile Ile Lys Lys Ile Arg Leu
        5
Ile Lys Lys
<210> 569
<211> 19
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
<400> 569
Lys Ile Gly Trp Lys Leu Arg Val Arg Ile Ile Arg Val Lys Ile Gly
                                                       15
                                   10
Arg Leu Arg
<210> 570
<211> 25
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE .
<400> 570
Lys Ile Val Ile Arg Ile Arg Ile Arg Leu Ile Arg Ile Arg Ile Arg
```

5

1

10

15

Lys Ile Val Lys Val Lys Arg Ile Arg
20 25

<210> 571

<211> 26

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 571

Arg Phe Ala Val Lys Ile Arg Leu Arg Ile Ile Lys Lys Ile Arg Leu

1 5 10 15

Ile Lys Lys Ile Arg Lys Arg Val Ile Lys
20 25

<210> 572

<211> 30

<212> PRT

<213> Artificial Sequence

<220>

<400> 572

Lys Ala Gly Trp Lys Leu Arg Val Arg Ile Ile Arg Val Lys Ile Gly
1 5 10 15

Arg Leu Arg Lys Ile Gly Trp Lys Lys Arg Val Arg Ile Lys 20 25 30

<210> 573

<211> 16

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: VIP MIMETIC

PEPTIDE

<400> 573

Arg Ile Tyr Val Lys Pro His Pro Arg Tyr Ile Lys Lys Ile Arg Leu 1 5 10 15

<210> 574

<211> 16

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PROTICE

<400> 574

Lys Pro Gly His Lys Ala Arg Pro His Ile Ile Arg Tyr Lys Ile Ile

1 5 10 15

<210> 575

<211> 19

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 575

Lys Ile Val Ile Arg Ile Arg Ile Arg Leu Ile Arg Ile Arg Ile Arg 1 5 10 15

Lys Ile Val

<210> 576

<211> 19

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: VIP MIMETIC

PEPTIDE

<400> 576

Arg Ile Ile Val Lys Ile Arg Leu Arg Ile Ile Lys Lys Ile Arg Leu

1 5 10 15

Ile Lys Lys

<210> 577

<211> 16

<212> PRT .

<213> Artificial Sequence

<220×

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 577

Arg Ile Tyr Val Ser Lys Ile Ser Ile Tyr Ile Lys Lys Ile Arg Leu

1 5 10 15

<210> 578

<211> 19

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 578

Lys Ile Val Ile Phe Thr Arg Ile Arg Leu Thr Ser Ile Arg Ile Arg 1 5 10 15

Ser Ile Val

<210> 579

<211> 16 ...

<212> PRT

<213> Artificial Sequence

<220> <223> Description of Artificial Sequence: VIP MIMETIC PEPTIDE <400> 579 Lys Pro Ile His Lys Ala Arg Pro Thr Ile Ile Arg Tyr Lys Met Ile 10 <210> 580 <211> 26 <212> PRT <213> Artificial Sequence <223> Description of Artificial Sequence: VIP MIMETIC PEPTIDE <220> <223> At position 1, disulfide bond to position 26 <220> <223> At position 26, disulfide bond to position 1 Xaa Cys Lys Gly Phe Phe Ala Leu Ile Pro Lys Ile Ile Ser Ser Pro 5 Leu Phe Lys Thr Leu Leu Ser Ala Val Cys 20 25 <210> 581 <211> 26 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE <400> 581

5

1

Cys Lys Lys Gly Phe Phe Ala Leu Ile Pro Lys Ile Ile Ser Ser_Pro

10

Leu Phe Lys Thr Leu Leu Ser Ala Val Cys
20 25

<210> 582

<211> 27

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 582

Cys Lys Lys Gly Phe Phe Ala Leu Ile Pro Lys Ile Ile Ser Ser
1 5 10 15

Pro Leu Phe Lys Thr Leu Leu Ser Ala Val Cys
20 25

<210> 583

<211> 17

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<220>

<223> At position 1, disulfide bond to position 17

<220>

<223> At position 17, disulfide bond to position 1

<400> 583

Xaa Cys Arg Ile Val Ile Arg Ile Arg Ile Arg Leu Ile Arg Ile Arg 1 1 5 15

Cys

<210> 584

```
<211> 19
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<220>
<223> At position 1, disulfide bond to position 19
<220>
<223> At position 19, disulfide bond to position 1
<400> 584
Xaa Cys Lys Pro Gly His Lys Ala Arg Pro His Ile Ile Arg Tyr Lys
                                    10
                  5
Ile Ile Cys
<210> 585
<211> 29
<212> PRT
<213> Artificial Sequence
<220>
```

PEPTIDE

<220>

<223> Description of Artificial Sequence:VIP MIMETIC

<223> At position 1, disulfide bond to position 29

<220>
<223> At position 29, disulfide bond to position 1

<400> 585

Xaa Cys Arg Phe Ala Val Lys Ile Arg Leu Arg Ile Ile Lys Lys Ile 1 5 10 15

Arg Leu Ile Lys Lys Ile Arg Lys Arg Val Ile Lys Cys 20 25

<210> 586

```
<211> 13
 <212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
 <400> 586
 Lys Leu Leu Lys Leu Leu Lys Leu Lys Cys
                5
 <210> 587
 <211> 12
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
 <400> 587
 Lys Leu Leu Lys Leu Leu Lys Leu Lys
                 5
 <210> 588
<211> 13
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: VIP MIMETIC
       PEPTIDE
 <400> 588
 Lys Leu Leu Lys Leu Lys Leu Lys Leu Lys Cys
                 5
  1
```

<220>

<223> Description of Artificial Sequence:VIP MIMETIC
 PEPTIDE

<400> 589

Lys Leu Leu Lys Leu Leu Leu Lys Leu Leu Lys

1 5 10

<210> 590

<211> 28

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 590

His Ser Asp Ala Val Phe Tyr Asp Asn Tyr Thr Arg Leu Arg Lys Gln
1 5 10 15

Met Ala Val Lys Lys Tyr Leu Asn Ser Ile Leu Asn 20 25

<210> 591

<211> 31

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 591

Asn Leu Glu His Ser Asp Ala Val Phe Tyr Asp Asn Tyr Thr Arg Leu
1 5 .10 15

Arg Lys Gln Met Ala Val Lys Lys Tyr Leu Asn Ser Ile Leu Asn 20 25 30

<210> 592

```
<211> 4
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
<220>
<223> At position 1, Xaa is absent or is ala, val,
      ala-val, val-ala, L-lys, D-lys, ala-lys, val-lys,
      ala-val-lys, val-ala-lys, or an ornithinyl residue
<220>
<223> At position 2, Xaa is L-lys, D-lys or an
      ornithinyl residue
<220>
<223> At position 3, Xaa is L-tyr, D-tyr, phe, trp or a
      p-aminophenylalanyl residue
<220>
<223> At position 4, Xaa is a hydrophobic aliphatic
      amino acid residue (X5), X5-leu, X5-norleucyl,
      X5-D-ala, X5-asn-ser, X5-asn-ser-ile,
      X5-asn-ser-tyr, X5-asn-ser-ile-leu,
      X5-asn-ser-tyr-leu,
<220>
<223> or X5-asn-ser-tyr-leu-asn
<400> 592
Xaa Xaa Xaa Xaa
  1
<210> 593
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
```

<223> At position 1, Xaa is either absent, a hydrophobic

<220>

```
aliphatic residue (X5), X5-asn, tyr-X5, lys-X5,
lyx-S5-asn, lys-tyr-X5, lys-tyr-X5-as,
lys-lys-tyr-X5, lys-lys-tyr-X5-asn,
val-lys-lys-tyr-X5,
```

<220>

<223> val-ala-lys-lys-tyr-X5-asn, or
 ala-val-lys-lys-tyr-X5-asn

<220>

<223> At position 3, Xaa is ile or tyr

<400> 593

Xaa Ser Xaa Leu Asn

<210> 594

<211> 7

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<220>

<223> At positions 1 and 6, Xaa are cross-linked amino
 acid residues in which the sidechain linker group
 is (CH2)m-Z-(CH2)n wherein Z is -CONH-, -NHCO-,
 -S-S-, -S(CH2)tCO-NH or -NH-CO(CH2)tS-; m is 1 or
2

<220>

<223> when Z is -NH-CO- or -NH-CO(CH2)tS-; n is 1 or 2
 when Z is -NH-CO-, -S-S- or -NH-CO(CH2)tS, or n is
2, 3 or 4 when Z is -CONH- or -S(CH2)tCO-NH-

<220>

<223> At position 5, Xaa is a hydrophobic aliphatic amino acid residue

<220>

<223> At position 7, Xaa is a covalent bond or Asn, Ser, Ile, Tyr, Leu, Asn-Ser, Asn-Ser-Ile, Asn-Ser-Tyr, Asn-Ser-Ile-Leu, Asn-Ser-Tyr-Leu, Asn-Ser-Ile-Leu-Asn or Asn-Ser-Tyr-Leu-Asn

```
<400> 594
Xaa Lys Lys Tyr Xaa Xaa Xaa
                 5
<210> 595
<211> 4
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
     PEPTIDE
<400> 595
Lys Lys Tyr Leu
<210> 596
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
<400> 596
Asn Ser Ile Leu Asn
<210> 597
<211> 4
<212> PRT
<213> Artificial Sequence
<220>
 <223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 597 ...
```

Lys Lys Tyr Leu

1

```
<210> 598
<211> 4
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
     PEPTIDE
<220>
<223> At position 4, D amino acid residue
<400> 598
Lys Lys Tyr Ala
 1
<210> 599
<211> 6
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 599
Ala Val Lys Lys Tyr Leu
  1 5
<210> 600
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 600 ...
```

Asn Ser Ile Leu Asn

1 5

```
<210> 601
<211> 4
<212> PRT
<213> Artificial Sequence
<220>
 <223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 601
 Lys Lys Tyr Val
  1
 <210> 602
 <211> 4
<212> PRT
<213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
 <220>
 <223> At position 3, Xaa is a lauric acid residue
 <400> 602
 Ser Ile Xaa Asn
  1
 <210> 603
 <211> 5
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: VIP MIMETIC
       PEPTIDE
 <220>
 <223> At position 5, Xaa is a norleucyl residue
```

```
<400> 603
Lys Lys Tyr Leu Xaa
 1
<210> 604
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 604
Asn Ser Tyr Leu Asn
<210> 605
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 605
Asn Ser Ile Tyr Asn
  1
<210> 606
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 606
Lys Lys Tyr Leu Pro Pro Asn Ser Ile Leu Asn
```

1 5 10

<210> 607
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<220>
<223> At position 1, Xaa is a lauric acid residue
<400> 607
Xaa Lys Lys Tyr Leu
1 5

<210> 608 <211> 5 <212> PRT <213> Artificial Sequence

<220>

<220>
<223> At position 1, Xaa is a caproic acid residue

<400> 608

Xaa Lys Lys Tyr Leu

<210> 609 <211> 4 <212> PRT <213> Artificial Sequence

```
<220>
<223> At position 4, Xaa is a norleucyl residue
<400> 609
Lys Lys Tyr Xaa
 1
<210> 610
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
     PEPTIDE
<400> 610
Val Lys Lys Tyr Leu
<210> 611
<211> 6
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
<400> 611
Leu Asn Ser Ile Leu Asn
                5 ·
<210> 612
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
```

```
<400> 612
 Tyr Leu Asn Ser Ile Leu Asn
 <210> 613
 <211> 5
 <212> PRT
  <213> Artificial Sequence
  <220>
  <223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
  <400> 613
  Lys Lys Tyr Leu Asn
   1
  <210> 614
  <211> 6
  <212> PRT
  <213> Artificial Sequence
  <220>
  <223> Description of Artificial Sequence: VIP MIMETIC
        PEPTIDE
  <400> 614
  Lys Lys Tyr Leu Asn Ser
                    5
  1
  <210> 615
  <211> 7
  <212> PRT
  <213> Artificial Sequence
  <220>
· <223> Description of Artificial Sequence: VIP MIMETIC
  <400> 615
```

Lys Lys Tyr Leu Asn Ser Ile

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5 1

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```
<210> 616
<211> 8
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
<400> 616
Lys Lys Tyr Leu Asn Ser Ile Leu
 1
<210> 617
<211> 4
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
<400> 617
Lys Lys Tyr Leu
  1
<210> 618
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
 <400> 618
Lys Lys Tyr Asp Ala
  1
```

```
<210> 619
<211> 6
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 619
Ala Val Lys Lys Tyr Leu
1
                5
<210> 620
<211> 5
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
<400> 620
Asn Ser Ile Leu Asn
<210> 621
<211> 4
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
<400> 621
Lys Lys Tyr Val
 1
```

<210> 622 <211> 4

```
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
<220>
<223> At position 3, Xaa is a lauric acid residue
<400> 622
Ser Ile Xaa Asn
<210> 623
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
<400> 623
Asn Ser Tyr Leu Asn
<210> 624
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 624
Asn Ser Ile Tyr Asn
```

<210> 625 <211> 5

```
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
   PEPTIDE
<220>
<223> At position 5, Xaa is a norleucyl residue
<400> 625
Lys Lys Tyr Leu Xaa
 1
<210> 626
<211> 11
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 626
Lys Lys Tyr Leu Pro Pro Asn Ser Ile Leu Asn
<210> 627
<211> 4
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 627
```

<210> 628 <211> 5

Lys Lys Tyr Leu

```
<212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence:VIP MIMETIC
       PEPTIDE
 <400> 628
 Lys Lys Tyr Asp Ala
 <210> 629
 <211> 6
 <212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence:VIP MIMETIC
       PEPTIDE
  <400> 629
  Ala Val Lys Lys Tyr Leu
  1
  <210> 630
  <211> 5
<212> PRT
  <213> Artificial Sequence
  <223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
  <400> 630
  Asn Ser Ile Leu Asn
· <210> 631
```

<211> 4 <212> PRT

<213> Artificial Sequence

```
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 631
Lys Lys Tyr Val
<210> 632
<211> 4
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
PEPTIDE
<220>
<223> At position 3, Xaa is a lauric acid residue
<400> 632
Ser Ile Xaa Asn
 1
<210> 633
<211> 6
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
<400> 633
Leu Ala Lys Lys Tyr Leu
                 5
 1
 <210> 634
 <211> 7
 <212> PRT--
 <213> Artificial Sequence
```

```
<220>
 <223> Description of Artificial Sequence: VIP MIMETIC
       PEPTIDE
 <400> 634
 Cys Ala Pro Lys Lys Tyr Leu
 <210> 635
<211> 4
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: VIP MIMETIC
       PEPTIDE
 <220>
 <223> At position 4, Xaa is a norleucyl residue
 <400> 635
 Lys Lys Tyr Xaa
  1
 <210> 636
 <211> 5
<212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: VIP MIMETIC
       PEPTIDE
 <400> 636
 Val Lys Lys Tyr Leu
  1
 <210> 637
 <211> 6
 <212> PRT ...
 <213> Artificial Sequence
```

```
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
<400> 637
Leu Asn Ser Ile Leu Asn
<210> 638
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
<400> 638
Tyr Leu Asn Ser Ile Leu Asn
                 5
 1
<210> 639
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
 <220>
<223> At position 5, Xaa is a norleucyl residue
 <400> 639
 Lys Lys Tyr Leu Xaa
  1
 <210> 640
 <211> 5
 <212> PRT ...
 <213> Artificial Sequence
```

```
<220>
 <223> Description of Artificial Sequence: VIP MIMETIC
       PEPTIDE
 <400> 640
 Lys Lys Tyr Leu Asn
 <210> 641
 <211> 6
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:VIP MIMETIC
 <400> 641
 Lys Lys Tyr Leu Asn Ser
  1
 <210> 642
 <211> 7
 <212> PRT
 <213> Artificial Sequence
<220>
 <223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE
 <400> 642
 Lys Lys Tyr Leu Asn Ser Ile
  1
              5 ·
 <210> 643
 <211> 8
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: VIP MIMETIC
```

PEPTIDE

```
<400> 643
 Lys Lys Tyr Leu Asn Ser Ile Leu
                 5
 <210> 644
 <211> 6
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:VIP MIMETIC
       PEPTIDE
 <400> 644
 Lys Lys Lys Tyr Leu Asp
<210> 645
 <211> 7
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:VIP MIMETIC
      PEPTIDE
 <220>
 <223> At positions 1, 6 disulfide cross-linked
 <400> 645
 Xaa Cys Lys Lys Tyr Leu Cys
  1
 <210> 646
 <211> 6
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: VIP MIMETIC
       PEPTIDE
```

```
<220>
<223> At positions 1, 6 cross-linked by S-CH2-CO
<400> 646
Cys Lys Lys Tyr Leu Lys
<210> 647
<211> 4
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
     PEPTIDE
<220>
<223> At position 4, D amino acid residue
<400> 647
Lys Lys Tyr Ala
 1 .
<210> 648
<211> 8
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VIP MIMETIC
      PEPTIDE .
<400> 648
Trp Trp Thr Asp Thr Gly Leu Trp
 1
<210> 649
<211> 8
<212> PRT-
<213> Artificial Sequence
```

```
<220>
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Trp Trp Asp Thr Arg Gly Leu Trp Val Trp Thr Ile
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Phe Trp Gly Asn Asp Gly Ile Trp Leu Glu Ser Gly
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PEPTIDE

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Asp Trp Asp Gln Phe Gly Leu Trp Arg Gly Ala Ala
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Arg Trp Asp Asp Asn Gly Leu Trp Val Val Leu
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Ser Gly Met Trp Ser His Tyr Gly Ile Trp Met Gly
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Gly Gly Arg Trp Asp Gln Ala Gly Leu Trp Val Ala

1 5 10

<210> 656

<211> 12

<212> PRT

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<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 656

Lys Leu Trp Ser Glu Gln Gly Ile Trp Met Gly Glu

1 5 10

<210> 657

<211> 10

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC
 PEPTIDE

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Cys Trp Ser Met His Gly Leu Trp Leu Cys
1 5 10

<210> 658

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:VIP MIMETIC PEPTIDE

<400> 658

Gly Cys Trp Asp Asn Thr Gly Ile Trp Val Pro Cys
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      PEPTIDE
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Asp Trp Asp Thr Arg Gly Leu Trp Val Tyr
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Ser Leu Trp Asp Glu Asn Gly Ala Trp Ile
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Lys Trp Asp Asp Arg Gly Leu Trp Met His
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        PEPTIDE
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<223> Description of Artificial Sequence:VIP MIMETIC

<213> Artificial Sequence

PEPTIDE

<220>

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Glu Trp Thr Asp Asn Gly Leu Trp Ala Leu
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Ser Trp Asp Glu Lys Gly Leu Trp Ser Ala
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 Ser His Leu Tyr Trp Gln Pro Tyr Ser Val Gln
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1 5 10

<210> 672
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Thr Leu Val Tyr Trp Gln Pro Tyr Ser Leu Gln Thr
1 5 10

<210> 673 <211> 12 <212> PRT <213> Artificial Sequence

<400> 673 Arg Gly Asp Tyr Trp Gln Pro Tyr Ser Val Gln Ser 1 5 10

<210> 674 <211> 12 <212> PRT <213> Artificial Sequence

<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST
PEPTIDE

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Val His Val Tyr Trp Gln Pro Tyr Ser Val Gln Thr
1 5 10

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Arg Leu Val Tyr Trp Gln Pro Tyr Ser Val Gln Thr
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Ser Arg Val Trp Phe Gln Pro Tyr Ser Leu Gln Ser
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<211> 12
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 Asn Met Val Tyr Trp Gln Pro Tyr Ser Ile Gln Thr
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<210> 678 <211> 12

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 Ser Val Val Phe Trp Gln Pro Tyr Ser Val Gln Thr
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 Thr Phe Val Tyr Trp Gln Pro Tyr Ala Leu Pro Leu
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Arg Leu Val Tyr Trp Gln Pro Tyr Ser Val Gln Arg
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Ser Pro Val Phe Trp Gln Pro Tyr Ser Ile Gln Ile
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Trp Ile Glu Trp Trp Gln Pro Tyr Ser Val Gln Ser
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<223> Description of Artificial Sequence:IL-1 ANTAGONIST

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  <211> 12
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        PEPTIDE
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Met Arg Val Phe Trp Gln Pro Tyr Ser Val Gln Asn

<400> 687

1 5 10

<210> 688 <211> 12' <212> PRT

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<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

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Lys Ile Val Tyr Trp Gln Pro Tyr Ser Val Gln Thr 1 5 10

<210> 689

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<400> 689

Arg His Leu Tyr Trp Gln Pro Tyr Ser Val Gln Arg
1 5 10

<210> 690

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<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

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Ala Leu Val Trp Trp Gln Pro Tyr Ser Glu Gln Ile 1 ... 5 10

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Ser Arg Val Trp Phe Gln Pro Tyr Ser Leu Gln Ser
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Trp Glu Gln Pro Tyr Ala Leu Pro Leu Glu
                 5
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<211> 12
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
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Gln Leu Val Trp Trp Gln Pro Tyr Ser Val Gln Arg
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<210> 694 <211> 12

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 Asp Leu Arg Tyr Trp Gln Pro Tyr Ser Val Gln Val
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 Glu Leu Val Trp Trp Gln Pro Tyr Ser Leu Gln Leu
                  5
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       PEPTIDE
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 Asp Leu Val Trp Trp Gln Pro Tyr Ser Val Gln Trp
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<210> 697 <211> 12 <212> PRT---<213> Artificial Sequence

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Asn Gly Asn Tyr Trp Gln Pro Tyr Ser Phe Gln Val
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Glu Leu Val Tyr Trp Gln Pro Tyr Ser Ile Gln Arg
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<211> 12
<212> PRT
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Glu Leu Met Tyr Trp Gln Pro Tyr Ser Val Gln Glu
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<210> 700
<211> 12
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST

PEPTIDE

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Asn Leu Leu Tyr Trp Gln Pro Tyr Ser Met Gln Asp
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<211> 12
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Gly Tyr Glu Trp Tyr Gln Pro Tyr Ser Val Gln Arg
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<211> 12
<212> PRT
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Ser Arg Val Trp Tyr Gln Pro Tyr Ser Val Gln Arg
1 5
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      PEPTIDE
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<400> 703

1 5 10

<210> 704 <211> 12 <212> PRT <213> Artificial Sequence

<400> 704
Gly Gly Gly Trp Trp Gln Pro Tyr Ser Val Gln Arg
1 5 10

<210> 705 <211> 12

<212> PRT

<213> Artificial Sequence

<2205

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 705
Val Gly Arg Trp Tyr Gln Pro Tyr Ser Val Gln Arg
1 5 10

<210> 706 <211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST
 PEPTIDE

<400> 706

Val His Val Tyr Trp Gln Pro Tyr Ser Val Gln Arg
1 5 10

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<210> 707
<211> 12
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      PEPTIDE
<400> 707
Gln Ala Arg Trp Tyr Gln Pro Tyr Ser Val Gln Arg
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<210> 708
<211> 12
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      PEPTIDE
<400> 708
Val His Val Tyr Trp Gln Pro Tyr Ser Val Gln Thr
                                     10
                5
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<211> 12
<212> PRT
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      PEPTIDE
<400> 709
Arg Ser Val Tyr Trp Gln Pro Tyr Ser Val Gln Arg
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<210> 710 <211> 12

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<212> PRT
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Thr Arg Val Trp Phe Gln Pro Tyr Ser Val Gln Arg
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<210> 711
<211> 12
<212> PRT
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<400> 711
Gly Arg Ile Trp Phe Gln Pro Tyr Ser Val Gln Arg
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<210> 712
<211> 12
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<210> 713 <211> 12

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Gly Arg Val Trp Phe Gln Pro Tyr Ser Val Gln Arg

<213> Artificial Sequence

<212> PRT-

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 Ala Arg Thr Trp Tyr Gln Pro Tyr Ser Val Gln Arg
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 <400> 714
 Ala Arg Val Trp Trp Gln Pro Tyr Ser Val Gln Met
                                      10
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       PEPTIDE
 <400> 715
 Arg Leu Met Phe Tyr Gln Pro Tyr Ser Val Gln Arg
  1
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 <210> 716
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST

<220>

PEPTIDE

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Glu Ser Met Trp Tyr Gln Pro Tyr Ser Val Gln Arg
                                     10
                 5
<210> 717
<211> 12
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      PEPTIDE
<400> 717
His Phe Gly Trp Trp Gln Pro Tyr Ser Val His Met
                 5
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<210> 718
<211> 12
<212> PRT
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<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 718
Ala Arg Phe Trp Trp Gln Pro Tyr Ser Val Gln Arg
                  5
<210> 719
<211> 12
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      PEPTIDE
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Arg Leu Val Tyr Trp Gln Pro Tyr Ala Pro Ile Tyr

<400> 719

1 5 10

<210> 720

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 720

Arg Leu Val Tyr Trp Gln Pro Tyr Ser Tyr Gln Thr 1 5 10

<210> 721

<211> 12

<212> PRT

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<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 721

Arg Leu Val Tyr Trp Gln Pro Tyr Ser Leu Pro Ile 1 5 10

<210> 722

<211> 12

<212> PRT

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<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 722

Arg Leu Val Tyr Trp Gln Pro Tyr Ser Val Gln Ala

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                                     10
                 5
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    PEPTIDE
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Ser Arg Val Trp Tyr Gln Pro Tyr Ala Gln Gly Leu
                 5
<210> 725
<211> 12
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      PEPTIDE
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Ser Arg Val Trp Tyr Gln Pro Tyr Ala Met Pro Leu
                  5
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<210> 726 <211> 12

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       PEPTIDE
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                  5
                                     10
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       PEPTIDE
<400> 727
 Ser Arg Val Trp Tyr Gln Pro Tyr Ser Leu Gly Leu
                  5
 <210> 728
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       PEPTIDE .
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<210> 729 <211> 12 <212> PRT ... <213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST
 PEPTIDE

<400> 729

Ser Arg Val Trp Tyr Gln Pro Tyr Ser Arg Gln Pro

1 5 10

<210> 730

<211> 12

<212> PRT

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<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 730

Ser Arg Val Trp Tyr Gln Pro Tyr Phe Val Gln Pro 1 5 10

<210> 731

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 731

Glu Tyr Glu Trp Tyr Gln Pro Tyr Ala Leu Pro Leu 1 5 10

<210> 732

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

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                 5
                                   10
<210> 733
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<212> PRT
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<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
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Ser Arg Ile Trp Trp Gln Pro Tyr Ala Leu Pro Leu
                5
<210> 734
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
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Asp Pro Leu Phe Trp Gln Pro Tyr Ala Leu Pro Leu
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<210> 735
<211> 12
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
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<400> 735 Ser Arg Gln Trp Val Gln Pro Tyr Ala Leu Pro Leu

1 5 10

<210> 736

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 736

Ile Arg Ser Trp Trp Gln Pro Tyr Ala Leu Pro Leu
1 5 10

<210> 737

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<212> PRT

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<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 737

Arg Gly Tyr Trp Gln Pro Tyr Ala Leu Pro Leu 1 5 10

<210> 738

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 738

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Arg Leu Leu Trp Val Gln Pro Tyr Ala Leu Pro Leu

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 Asp Ala Tyr Trp Val Gln Pro Tyr Ala Leu Pro Leu
                                     10
                 5
 <210> 741
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       PEPTIDE
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 Trp Ser Gly Tyr Phe Gln Pro Tyr Ala Leu Pro Leu
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                   5
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<210> 742 <211> 12

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Asn Ile Glu Phe Trp Gln Pro Tyr Ala Leu Pro Leu
                 5
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<400> 743
Thr Arg Asp Trp Val Gln Pro Tyr Ala Leu Pro Leu
          5
<210> 744
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      PEPTIDE
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Asp Ser Ser Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
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<210> 745 <211> 12 <212> PRT ---<213> Artificial Sequence

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<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 745

Ile Gly Asn Trp Tyr Gln Pro Tyr Ala Leu Pro Leu

1 5 10
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<210> 746
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<400> 746

1 5 10

Asn Leu Arg Trp Asp Gln Pro Tyr Ala Leu Pro Leu

<210> 747 <211> 12 <212> PRT <213> Artificial Sequence <220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 747
Leu Pro Glu Phe Trp Gln Pro Tyr Ala Leu Pro Leu
1 5 10

<210> 748
<211> 12
<212> PRT
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<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST
PEPTIDE

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Asp Ser Tyr Trp Trp Gln Pro Tyr Ala Leu Pro Leu
  1
                5
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<210> 749
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<212> PRT
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      PEPTIDE
<400> 749
Arg Ser Gln Tyr Tyr Gln Pro Tyr Ala Leu Pro Leu
                5
<210> 750
<211> 12
<212> PRT
<213> Artificial Sequence
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      PEPTIDE
<400> 750
Ala Arg Phe Trp Leu Gln Pro Tyr Ala Leu Pro Leu
                 5
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<210> 751
<211> 12
<212> PRT
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<220>
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Asn Ser Tyr Phe Trp Gln Pro Tyr Ala Leu Pro Leu

PEPTIDE

<400> 751

1 5 10

<210> 752
<211> 12
<212> PRT
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<400> 752
Arg Phe Met Tyr Trp Gln Pro Tyr Ser Val Gln Arg

1 5 10

<210> 753 <211> 12 <212> PRT <213> Artificial Sequence

PEPTIDE

<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST

<400> 753
Ala His Leu Phe Trp Gln Pro Tyr Ser Val Gln Arg
1 5 10

<210> 754 <211> 9 <212> PRT <213> Artificial Sequence <220>

<400> 754
Trp Trp Gln Pro Tyr Ala Leu Pro Leu
1 ... 5

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      PEPTIDE
<400> 755
Tyr Tyr Gln Pro Tyr Ala Leu Pro Leu
                5
<210> 756
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<212> PRT
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      PEPTIDE
<400> 756
Tyr Phe Gln Pro Tyr Ala Leu Gly Leu
                5
<210> 757
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 757
Tyr Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
                 5
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<210> 758 <211> 10

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<212> PRT
<213> Artificial Sequence
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<400> 758
 Arg Trp Trp Gln Pro Tyr Ala Thr Pro Leu
 <210> 759
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 <400> 759
 Gly Trp Tyr Gln Pro Tyr Ala Leu Gly Phe
       5
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 <211> 10
<212> PRT
 <213> Artificial Sequence
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       PEPTIDE
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 Tyr Trp Tyr Gln Pro Tyr Ala Leu Gly Leu
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<212> PRT-

<213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
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Ile Trp Tyr Gln Pro Tyr Ala Met Pro Leu
                  5
<210> 762
<211> 10
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      PEPTIDE
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Ser Asn Met Gln Pro Tyr Gln Arg Leu Ser
                 5
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<210> 763
<211> 20
<212> PRT
<213> Artificial Sequence
<220>
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      PEPTIDE
<400> 763
Thr Phe Val Tyr Trp Gln Pro Tyr Ala Val Gly Leu Pro Ala Ala Glu
                                    10
                  5
  1
Thr Ala Cys Asn
              20
<210> 764
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 <212> PRT ...
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<213> Artificial Sequence

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<220>
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      PEPTIDE
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Thr Phe Val Tyr Trp Gln Pro Tyr Ser Val Gln Met Thr Ile Thr Gly
                 5
                                   10
Lys Val Thr Met
             20
<210> 765
<211> 20
<212> PRT
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<220>
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      PEPTIDE
<400> 765
Thr Phe Val Tyr Trp Gln Pro Tyr Ser Ser His Xaa Xaa Val Pro Xaa
                                   10
Gly Phe Pro Leu
<210> 766
<211> 20
 <212> PRT
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      PEPTIDE
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 Thr Phe Val Tyr Trp Gln Pro Tyr Tyr Gly Asn Pro Gln Trp Ala Ile
                                   10
                 5
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His Val Arg His

... 20

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<210> 767
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      PEPTIDE
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Thr Phe Val Tyr Trp Gln Pro Tyr Val Leu Leu Glu Leu Pro Glu Gly
                  5
Ala Val Arg Ala
<210> 768
<211> 20
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      PEPTIDE
<400> 768
Thr Phe Val Tyr Trp Gln Pro Tyr Val Asp Tyr Val Trp Pro Ile Pro
  1
                                      10
Ile Ala Gln Val
             20
<210> 769
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<212> PRT
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 769
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Gly Trp Tyr Gln Pro Tyr Val Asp Gly Trp Arg

5 10

<210> 770

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<400> 770 ·

Arg Trp Glu Gln Pro Tyr Val Lys Asp Gly Trp Ser

<210> 771

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST
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<400> 771

Glu Trp Tyr Gln Pro Tyr Ala Leu Gly Trp Ala Arg

1 5 10

<210> 772

<211> 10

<212> PRT

<213> Artificial Sequence

<220>

<400> 772

Gly Trp Trp Gln Pro Tyr Ala Arg Gly Leu
1 5 10

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Leu Phe Glu Gln Pro Tyr Ala Lys Ala Leu Gly Leu
                  5
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      PEPTIDE
<400> 774
Gly Trp Glu Gln Pro Tyr Ala Arg Gly Leu Ala Gly
                 5
<210> 775
<211> 12
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 775
Ala Trp Val Gln Pro Tyr Ala Thr Pro Leu Asp Glu
                 5
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<210> 776 <211> 12

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      PEPTIDE
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Met Trp Tyr Gln Pro Tyr Ser Ser Gln Pro Ala Glu
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Gly Trp Thr Gln Pro Tyr Ser Gln Gln Gly Glu Val
                  5
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 Asp Trp Phe Gln Pro Tyr Ser Ile Gln Ser Asp Glu
                   5
   1
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290

<210> 779
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      PEPTIDE
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Pro Trp Ile Gln Pro Tyr Ala Arg Gly Phe Gly
                 5
<210> 780
<211> 12
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Arg Pro Leu Tyr Trp Gln Pro Tyr Ser Val Gln Val
          5
 1
<210> 781
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    PEPTIDE
<400> 781
Thr Leu Ile Tyr Trp Gln Pro Tyr Ser Val Gln Ile
                 5
                                     10
<210> 782
<211> 12
<212> PRT
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<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
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PEPTIDE

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Arg Phe Asp Tyr Trp Gln Pro Tyr Ser Asp Gln Thr
 1
                 5
<210> 783
<211> 12
<212> PRT
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<220>
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      PEPTIDE
<400> 783
Trp His Gln Phe Val Gln Pro Tyr Ala Leu Pro Leu
                5
<210> 784
<211> 17
<212> PRT
<213> Artificial Sequence
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     PEPTIDE
<400> 784
Glu Trp Asp Ser Val Tyr Trp Gln Pro Tyr Ser Val Gln Thr Leu Leu
                  5
                                   10
Arg
<210> 785
<211> 17
<212> PRT
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
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PEPTIDE

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Trp Glu Gln Asn Val Tyr Trp Gln Pro Tyr Ser Val Gln Ser Phe Ala
                                   10
 1
                  5
Asp
<210> 786
<211> 16
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<220>
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     PEPTIDE
<400> 786
Ser Asp Val Val Tyr Trp Gln Pro Tyr Ser Val Gln Ser Leu Glu Met
                                    10
                  5
<210> 787
<211> 17
<212> PRT
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      PEPTIDE
<400> 787
Tyr Tyr Asp Gly Val Tyr Trp Gln Pro Tyr Ser Val Gln Val Met Pro
                                     10
 1
                  5
Ala
<210> 788
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<212> PRT

<213> Artificial Sequence

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Ser Asp Ile Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
<210> 789
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 Gln Arg Ile Trp Trp Gln Pro Tyr Ala Leu Pro Leu
                  5 .
 <210> 790
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 <400> 790
 Ser Arg Ile Trp Trp Gln Pro Tyr Ala Leu Pro Leu
                                     10
                  5
  <210> 791
  <211> 12
  <212> PRT
  <213> Artificial Sequence
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST

<220> ---

PEPTIDE

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Arg Ser Leu Tyr Trp Gln Pro Tyr Ala Leu Pro Leu
                  5
                                     10
 1
<210> 792
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<212> PRT
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      PEPTIDE
<400> 792
Thr Ile Ile Trp Glu Gln Pro Tyr Ala Leu Pro Leu
                 5
<210> 793
<211> 12
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<220>
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      PEPTIDE
<400> 793
Trp Glu Thr Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
                 5
  1
<210> 794
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<212> PRT
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      PEPTIDE
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Ser Tyr Asp Trp Glu Gln Pro Tyr Ala Leu Pro Leu

1 5 10

<210> 795

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 795

Ser Arg Ile Trp Cys Gln Pro Tyr Ala Leu Pro Leu 1 5 10

<210> 796

<211> 12

<212> PRT

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<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 796

Glu Ile Met Phe Trp Gln Pro Tyr Ala Leu Pro Leu
1 5 10

<210> 797

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 797

Asp Tyr Val Trp Gln Gln Pro Tyr Ala Leu Pro Leu

1 ... 5 .

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<210> 798
<211> 15
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<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 798
Met Asp Leu Leu Val Gln Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
                 5
                                    10
<210> 799
<211> 15
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST
      PEPTIDE
<400> 799
Gly Ser Lys Val Ile Leu Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
                           10
                5
<210> 800
<211> 15
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<400> 800
Arg Gln Gly Ala Asn Ile Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
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<210> 801 . <211> 15

<212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence:IL-1 ANTAGONIST <400> 801 Gly Gly Gly Asp Glu Pro Trp Tyr Gln Pro Tyr Ala Leu Pro Leu 10 5 <210> 802 . <211> 15 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE <400> 802 Ser Gln Leu Glu Arg Thr Trp Tyr Gln Pro Tyr Ala Leu Pro Leu 10 5 <210> 803 <211> 15 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE <400> 803 Glu Thr Trp Val Arg Glu Trp Tyr Gln Pro Tyr Ala Leu Pro Leu 10

<210> 804 <211> 15 <212> PRT <213> Artificial Sequence <220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST
 PEPTIDE

<400> 804

Lys Lys Gly Ser Thr Gln Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
1 5 10 15

<210> 805

<211> 15

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 805

Leu Gln Ala Arg Met Asn Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
1 5 10 15

<210> 806

<211> 15

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 806

Glu Pro Arg Ser Gln Lys Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
1 5 10 15

<210> 807

<211> 15

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 807 Val Lys Gln Lys Trp Arg Trp Tyr Gln Pro Tyr Ala Leu Pro Leu 5 10 <210> 808 <211> 15 . <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE <400> 808 Leu Arg Arg His Asp Val Trp Tyr Gln Pro Tyr Ala Leu Pro Leu 10 5 <210> 809 <211> 15 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE <400> 809 Arg Ser Thr Ala Ser Ile Trp Tyr Gln Pro Tyr Ala Leu Pro Leu 10 5 <210> 810 <211> 15 <212> PRT <213> Artificial Sequence

<220>

<400> 810

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Glu Ser Lys Glu Asp Gln Trp Tyr Gln Pro Tyr Ala Leu Pro Leu

1 5 10 15

<210> 811

<211> 15

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 811

Glu Gly Ser Arg Glu Gly Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
1 5 10 15

<210> 812

<211> 15

<212> PRT

<213> Artificial Sequence

-220-

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 812

Glu Gly Ser Arg Glu Gly Trp Tyr Gln Pro Tyr Ala Leu Pro Leu 1 5 10 15

<210> 813

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<400> 813

Val Ile Glu Trp Trp Gln Pro Tyr Ala Leu Pro Leu
1 5 10

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<210> 814
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<212> PRT
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<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<400> 814
Val Trp Tyr Trp Glu Gln Pro Tyr Ala Leu Pro Leu
                5
<210> 815
<211> 12
<212> PRT
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      PEPTIDE
<400> 815
Ala Ser Glu Trp Trp Gln Pro Tyr Ala Leu Pro Leu
                  5
  1
<210> 816
<211> 12
<212> PRT
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      PEPTIDE
 <400> 816
 Phe Tyr Glu Trp Trp Gln Pro Tyr Ala Leu Pro Leu
                   5
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<210> 817 <211> 12

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<212> PRT
<213> Artificial Sequence
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 817
Glu Gly Trp Trp Val Gln Pro Tyr Ala Leu Pro Leu
<210> 818
<211> 12
<212> PRT
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      PEPTIDE
<400> 818
Trp Gly Glu Trp Leu Gln Pro Tyr Ala Leu Pro Leu
<210> 819
<211> 12
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       PEPTIDE
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Asp Tyr Val Trp Glu Gln Pro Tyr Ala Leu Pro Leu
                   5
 <210> 820
 <211> 12
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<212> PRT

<213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 820
Ala His Thr Trp Trp Gln Pro Tyr Ala Leu Pro Leu
                 5
<210> 821
<211> 12
<212> PRT
<213> Artificial Sequence
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
   PEPTIDE
<400> 821
 Phe Ile Glu Trp Phe Gln Pro Tyr Ala Leu Pro Leu
                 5
 <210> 822
 <211> 12
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <400> 822
 Trp Leu Ala Trp Glu Gln Pro Tyr Ala Leu Pro Leu
             5
  1
  <210> 823
  <211> 12
  <212> PRT
  <213> Artificial Sequence
  <220>
  <223> Description of Artificial Sequence: IL-1 ANTAGONIST
        PEPTIDE
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<400> 823 Val Met Glu Trp Trp Gln Pro Tyr Ala Leu Pro Leu 5 <210> 824 <211> 11 <212> PRT <213> Artificial Sequence <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE <400> 824 Glu Arg Met Trp Gln Pro Tyr Ala Leu Pro Leu <210> 825 <211> 12 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE <400> 825 Asn Xaa Xaa Trp Xaa Xaa Pro Tyr Ala Leu Pro Leu 10 5 <210> 826 <211> 12 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

Trp Gly Asn Trp Tyr Gln Pro Tyr Ala Leu Pro Leu

<400> 826

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10 5 1

<210> 827

<211> 12

<212> PRT

<213> Artificial Sequence

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 827

Thr Leu Tyr Trp Glu Gln Pro Tyr Ala Leu Pro Leu 5 . 1

<210> 828

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 828

Val Trp Arg Trp Glu Gln Pro Tyr Ala Leu Pro Leu 10 5

<210> 829

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 829

Leu Leu Trp Thr Gln Pro Tyr Ala Leu Pro Leu 10 5

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<210> 830
<211> 12
<212> PRT
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 830
Ser Arg Ile Trp Xaa Xaa Pro Tyr Ala Leu Pro Leu
                5
<210> 831
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 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
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 Ser Asp Ile Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
  1 5
 <210> 832
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       PEPTIDE
  <400> 832
  Trp Gly Tyr Tyr Xaa Xaa Pro Tyr Ala Leu Pro Leu
                    5
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<210> 833 <211> 12

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<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 833
Thr Ser Gly Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
<210> 834
<211> 12
<212> PRT
<213> Artificial Sequence
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      PEPTIDE
<400> 834
Val His Pro Tyr Xaa Xaa Pro Tyr Ala Leu Pro Leu
                  5
 <210> 835
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       PEPTIDE
 <400> 835
 Glu His Ser Tyr Phe Gln Pro Tyr Ala Leu Pro Leu
                 5
 <210> 836
 <211> 12
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<212> PRT

<213> Artificial Sequence

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<220>
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     PEPTIDE
<400> 836
Xaa Xaa Ile Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
 1 5
<210> 837
<211> 12
<212> PRT
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 837
Ala Gln Leu His Ser Gln Pro Tyr Ala Leu Pro Leu
 1
                5
<210> 838
<211> 12
<212> PRT
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<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 838
Trp Ala Asn Trp Phe Gln Pro Tyr Ala Leu Pro Leu
 1
                 5
                        10
<210> 839
<211> 12
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
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PEPTIDE

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<400> 839
Ser Arg Leu Tyr Ser Gln Pro Tyr Ala Leu Pro Leu
                5
<210> 840
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 840
Gly Val Thr Phe Ser Gln Pro Tyr Ala Leu Pro Leu
                5
<210> 841
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 841
Ser Ile Val Trp Ser Gln Pro Tyr Ala Leu Pro Leu
 1 . 5
<210> 842
 <211> 12
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <400> 842
 Ser Arg Asp Leu Val Gln Pro Tyr Ala Leu Pro Leu
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1 5 10

<210> 843

<211> 17

<212> PRT

<213> Artificial Sequence

<220>

<400> 843

His Trp Gly His Val Tyr Trp Gln Pro Tyr Ser Val Gln Asp Asp Leu

1 5 10 15

Gly

<210> 844

<211> 17

<212> PRT

<213> Artificial Sequence

<220>

<400> 844

Ser Trp His Ser Val Tyr Trp Gln Pro Tyr Ser Val Gln Ser Val Pro 1 5 10 15

Glu

<210> 845

<211> 17

<212> PRT

<213> Artificial Sequence

<220>

```
<400> 845
 Trp Arg Asp Ser Val Tyr Trp Gln Pro Tyr Ser Val Gln Pro Glu Ser
                 5
                                   10
                                           15
Ala
 <210> 846
 <211> 17
 <212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <400> 846
 Thr Trp Asp Ala Val Tyr Trp Gln Pro Tyr Ser Val Gln Lys Trp Leu
                                   10
                 5
 Asp
 <210> 847
 <211> 17
 <212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <400> 847
 Thr Pro Pro Trp Val Tyr Trp Gln Pro Tyr Ser Val Gln Ser Leu Asp
                                                      15
                          10
                   5
  Pro
```

<210> 848 <211> 17

```
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 848
Tyr Trp Ser Ser Val Tyr Trp Gln Pro Tyr Ser Val Gln Ser Val His
                  5
                                    10
Ser
<210> 849
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
<400> 849
Tyr Trp Tyr Gln Pro Tyr Ala Leu Gly Leu
            5
<210> 850
<211> 10
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <400> 850
 Tyr Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
                              : 10
                  5
  1
```

<210> 851 <211> 10

```
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 851
Glu Trp Ile Gln Pro Tyr Ala Thr Gly Leu
         5
<210> 852
<211> 10
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 852
Asn Trp Glu Gln Pro Tyr Ala Lys Pro Leu
          5
<210> 853
<211> 10
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <400> 853
 Ala Phe Tyr Gln Pro Tyr Ala Leu Pro Leu
                 5
 <210> 854
 <211> 10 ---
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<212> PRT

<213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 854
Phe Leu Tyr Gln Pro Tyr Ala Leu Pro Leu
        5
<210> 855
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 855
Val Cys Lys Gln Pro Tyr Leu Glu Trp Cys
        5
<210> 856
<211> 21
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <400> 856
Glu Thr Pro Phe Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro
                                    10
                  5
 Tyr Ala Leu Pro Leu
              20
 <210> 857
 <211> 21...
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<212> PRT

<213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 857
Gln Gly Trp Leu Thr Trp Gln Asp Ser Val Asp Met Tyr Trp Gln Pro
Tyr Ala Leu Pro Leu
             20
<210> 858
<211> 21
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 858
Phe Ser Glu Ala Gly Tyr Thr Trp Pro Glu Asn Thr Tyr Trp Gln Pro
                                      10
                  5
  1
Tyr Ala Leu Pro Leu
             20
<210> 859
<211> 21
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <400> 859
 Thr Glu Ser Pro Gly Gly Leu Asp Trp Ala Lys Ile Tyr Trp Gln Pro
                                      10
                   5
   1
```

Tyr Ala Leu Pro Leu

20

```
<210> 860
<211> 21
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST
<400> 860
Asp Gly Tyr Asp Arg Trp Arg Gln Ser Gly Glu Arg Tyr Trp Gln Pro
                                     10
                 5
Tyr Ala Leu Pro Leu
        20
<210> 861
<211> 21
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <400> 861
 Thr Ala Asn Val Ser Ser Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro
                                                       15
                                     10
 Tyr Ala Leu Pro Leu
            20
 <210> 862
 <211> 21
 <212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
```

Ser Val Gly Glu Asp His Asn Phe Trp Thr Ser Glu Tyr Trp Gln Pro

<400> 862

1 5 10 15

Tyr Ala Leu Pro Leu 20

<210> 863

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 863

Met Asn Asp Gln Thr Ser Glu Val Ser Thr Phe Pro Tyr Trp Gln Pro 1 5 10 15

Tyr Ala Leu Pro Leu 20

<210> 864

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<400> 864

Ser Trp Ser Glu Ala Phe Glu Gln Pro Arg Asn Leu Tyr Trp Gln Pro 1 5 10 15

Tyr Ala Leu Pro Leu

20

<210> 865

<211> 21 ...

<212> PRT

<213> Artificial Sequence

<220>

<400> 865

Gln Tyr Ala Glu Pro Ser Ala Leu Asn Asp Trp Gly Tyr Trp Gln Pro 1 5 10 15

Tyr Ala Leu Pro Leu 20

<210> 866

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<400> 866

Asn Gly Asp Trp Ala Thr Ala Asp Trp Ser Asn Tyr Tyr Trp Gln Pro

Tyr Ala Leu Pro Leu 20

<210> 867

<211> 15

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL·1 ANTAGONIST PEPTIDE

<400> 867

Thr His Asp Glu His Ile Tyr Trp Gln Pro Tyr Ala Leu Pro Leu
1 5 10 15

<210> 868

<211> 21

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<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 868
Met Leu Glu Lys Thr Tyr Thr Trp Thr Pro Gly Tyr Trp Gln Pro
                 5
                                   10
Tyr Ala Leu Pro Leu
            20
<210> 869
<211> 20
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
Trp Ser Asp Pro Leu Thr Arg Asp Ala Asp Leu Tyr Trp Gln Pro Tyr
                 5 10
Ala Leu Pro Leu
             20
<210> 870
<211> 21
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 870
Ser Asp Ala Phe Thr Thr Gln Asp Ser Gln Ala Met Tyr Trp Gln Pro
                                    10
                 5
```

Tyr Ala Leu Pro Leu

20

<210> 871

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<400> 871

Gly Asp Asp Ala Ala Trp Arg Thr Asp Ser Leu Thr Tyr Trp Gln Pro 1 5 10 15

Tyr Ala Leu Pro Leu 20

<210> 872

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 872

Ala Ile Ile Arg Gln Leu Tyr Arg Trp Ser Glu Met Tyr Trp Gln Pro 1 5 10 15

Tyr Ala Leu Pro Leu 20

<210> 873

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<400> 873

Glu Asn Thr Tyr Ser Pro Asn Trp Ala Asp Ser Met Tyr Trp Gln Pro 1 5 10 15

Tyr Ala Leu Pro Leu 20

<210> 874

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<400> 874

Met Asn Asp Gln Thr Ser Glu Val Ser Thr Phe Pro Tyr Trp Gln Pro 1 5 10 15

Tyr Ala Leu Pro Leu 20

<210> 875

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<400> 875

Ser Val Gly Glu Asp His Asn Phe Trp Thr Ser Glu Tyr Trp Gln Pro 1 5 10 15

Tyr Ala Leu Pro Leu

20

<210> 876

<211> 21

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<212> PRT
<213> Artificial Sequence
<220>
```

<223> Description of Artificial Sequence: IL-1 ANTAGONIST

<400> 876

Gln Thr Pro Phe Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro 1 5 10 15

Tyr Ala Leu Pro Leu 20

<210> 877 <211> 21 <212> PRT

<213> Artificial Sequence

<220>

<400> 877

Glu Asn Pro Phe Thr Trp Gln Glu Ser Asn Ala Tyr Tyr Trp Gln Pro 1 5 10 15

Tyr Ala Leu Pro Leu 20

<210> 878 <211> 21

<212> PRT

<213> Artificial Sequence

<220>

<400> 878

Val Thr Pro Phe Thr Trp Glu Asp Ser Asn Val Phe Tyr Trp Gln Pro

1 5 10 15

Tyr Ala Leu Pro Leu

PCT/US99/25044 WO 00/24782

20

<210> 879

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST

<400> 879

Gln Ile Pro Phe Thr Trp Glu Gln Ser Asn Ala Tyr Tyr Trp Gln Pro 10 5

Tyr Ala Leu Pro Leu 20

<210> 880

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 880

Gln Ala Pro Leu Thr Trp Gln Glu Ser Ala Ala Tyr Tyr Trp Gln Pro 10

Tyr Ala Leu Pro Leu 20

<210> 881

<211> 21 <212> PRT

<213> Artificial Sequence

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 881 Glu Pro Thr Phe Thr Trp Glu Glu Ser Lys Ala Thr Tyr Trp Gln Pro 5 10 Tyr Ala Leu Pro Leu 20 <210> 882 <211> 21 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE <400> 882 Thr Thr Thr Leu Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro 15 5 Tyr Ala Leu Pro Leu 20 <210> 883 <211> 21 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE <400> 883 Glu Ser Pro Leu Thr Trp Glu Glu Ser Ser Ala Leu Tyr Trp Gln Pro 15 10 5 1 Tyr Ala Leu Pro Leu

<210> 884 <211> 21 20

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<212> PRT
  <213> Artificial Sequence
  <220>
  <223> Description of Artificial Sequence: IL-1 ANTAGONIST
        PEPTIDE
  <400> 884
  Glu Thr Pro Leu Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro
                                      10
  Tyr Ala Leu Pro Leu
               20
  <210> 885
  <211> 21
  <212> PRT
<213> Artificial Sequence
  <220>
  <223> Description of Artificial Sequence: IL-1 ANTAGONIST
        PEPTIDE
  <400> 885
  Glu Ala Thr Phe Thr Trp Ala Glu Ser Asn Ala Tyr Tyr Trp Gln Pro
                    5
                                       10
  Tyr Ala Leu Pro Leu
               20
  <210> 886
   <211> 21
   <212> PRT
   <213> Artificial Sequence
   <223> Description of Artificial Sequence: IL-1 ANTAGONIST
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<400> 886

Glu Ala Leu Phe Thr Trp Lys Glu Ser Thr Ala Tyr Tyr Trp Gln Pro 15 ~ 10 5

Tyr Ala Leu Pro Leu

PEPTIDE

20

<210> 887

<211> 20

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 887

Ser Thr Pro Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro Tyr

1 5 10 15

Ala Leu Pro Leu

20

<210> 888

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<400> 888

Glu Thr Pro Phe Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro

Tyr Ala Leu Pro Leu

20

<210> 889

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 889 Lys Ala Pro Phe Thr Trp Glu Glu Ser Gln Ala Tyr Tyr Trp Gln Pro 10 Tyr Ala Leu Pro Leu 20 <210> 890 <211> 21 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE <400> 890 Ser Thr Ser Phe Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro 15 10 1 5 Tyr Ala Leu Pro Leu 20 <210> 891 <211> 21 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE <400> 891 Asp Ser Thr Phe Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro 15 10 5 1 Tyr Ala Leu Pro Leu

<210> 892 <211> 21 20

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<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<400> 892
Tyr Ile Pro Phe Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro
                5
                                 10
Tyr Ala Leu Pro Leu
            20
<210> 893
<211> 21
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 893
Gln Thr Ala Phe Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro
                                   10
Tyr Ala Leu Pro Leu
            20
<210> 894
<211> 21
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 894
Glu Thr Leu Phe Thr Trp Glu Glu Ser Asn Ala Thr Tyr Trp Gln Pro
 1 ··· 5 · 10 · 15 .
```

Tyr Ala Leu Pro Leu

20

```
<210> 895
<211> 21
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 895
Val Ser Ser Phe Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro
                                     10
Tyr Ala Leu Pro Leu
            20
<210> 896
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST
      PEPTIDE
<400> 896
Gln Pro Tyr Ala Leu Pro Leu
                 5
<210> 897
<211> 11
<212> PRT
<213> Artificial Sequence
 <220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <220>
 <223> At position 1, Xaa is a phosphotyrosyl residue
```

```
<220>
<223> At position 2, Xaa is a 1-napthylalanyl residue
<220>
<223> At position 6, Xaa is an azetidine residue
<400> 897
Xaa Xaa Pro Tyr Gln Xaa Tyr Ala Leu Pro Leu
                5
<210> 898
<211> 21
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
    PEPTIDE
<400> 898
Thr Ala Asn Val Ser Ser Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro
                 5
                                    10
Tyr Ala Leu Pro Leu
             20
<210> 899
<211> 15
<212> PRT
<213> Artificial Sequence
<220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <400> 899
 Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro Tyr Ala Leu Pro Leu
                                    10
  1
```

<210> 900 <211> 15

```
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 10, Xaa is an azetidine residue
<400> 900
Phe Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr Ala Leu Pro Leu
                                     10
                  5
<210> 901
<211> 15
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 10, Xaa is an azetidine residue
<400> 901
Phe Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Tyr Ala Leu Pro Leu
                                     10
                   5
<210> 902
<211> 21
<212> PRT
<213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <400> 902
 Glu Thr Pro Phe Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro
                                      10
                   5
  1
```

Tyr Ala Leu Pro Leu

20

<210> 904 <211> 16 <212> PRT <213> Artificial Sequence

<220> <223> Description of Artificial Sequence:IL-1 ANTAGONIST

PEPTIDE

<210> 905 <211> 17 <212> PRT <213> Artificial Sequence

<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

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<400> 905
Gly Asp Val Ala Glu Tyr Trp Gln Pro Tyr Ala Leu Pro Leu Thr Ser
                                    10
                5
Leu
<210> 906
<211> 18
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
Ser Trp Thr Asp Tyr Gly Tyr Trp Gln Pro Tyr Ala Leu Pro Ile Ser
                                                         15
                                    10
                 5
  1
Gly Leu
<210> 907
<211> 8
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 4, Xaa is prolyl or an azetidine
      residue
 <220>
 <223> At position 6, Xaa is S, A, V or L
 <400> 907
 Xaa Xaa Gln Xaa Tyr Xaa Xaa Xaa
                  5
```

```
<210> 908
<211> 8
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 1, Xaa is Y, W or F
<220>
<223> At position 4, Xaa is prolyl or an azetidine
      residue
<220>
<223> At position 6, Xaa is S, A, V or L
<400> 908
Xaa Xaa Gln Xaa Tyr Xaa Xaa Xaa
                 5
<210> 909
<211> 8
<212> PRT
<213> Artificial Sequence
<220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <220>
 <223> At position 1, Xaa is Y, W or F
 <223> At position 2, Xaa is E, F, V, W or Y
 <223> At position 4, Xaa is prolyl or an azetidine
       residue
 <220>
 <223> At position 6, Xaa is S, A, V or L
```

```
<220>
<223> At position 7, Xaa is M, F, V, R, Q, K, T, S, D,
     L, I or E
<220>
<223> At position 8, Xaa is E, L, W, V, H, I, G, A, D,
     L, Y, N, Q or P
<400> 909
Xaa Xaa Gln Xaa Tyr Xaa Xaa Xaa
                 5
 1
<210> 910
<211> 9
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 1, Xaa is V, L, I, E, P, G, Y, M, T or
      D
<220>
<223> At position 2, Xaa is Y, W or F
<220>
<223> At position 3, Xaa is E, F, V, W or Y
<220>
<223> At position 5, Xaa is prolyl or an azetidine
      residue
<220>
<223> At position 7, Xaa is S, A, V or L
<220>
<223> At position 8, Xaa is M, F, V, R, Q, K, T, S, D,
      L, I or E
<220>
```

<223> At position 9, Xaa is E, L, W, V, H, I, G, A, D,

L, Y, N, Q or P

```
<400> 910
Xaa Xaa Xaa Gln Xaa Tyr Xaa Xaa Xaa
                 5
<210> 911
<211> 15
<212> PRT
<213> Artificial Sequence
<220> .
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 911
Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro Tyr Ala Leu Pro Leu
                  5
<210> 912
<211> 15
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 10, Xaa is an azetidine residue
Phe Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr Ala Leu Pro Leu
                                     10
                   5
 <210> 913
 <211> 15
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
```

PEPTIDE

```
<400> 913
 Phe Glu Trp Thr Pro Gly Trp Tyr Gln Pro Tyr Ala Leu Pro Leu
  1 5 10 15
 <210> 914
 <211> 15
  <212> PRT
  <213> Artificial Sequence
 <220>
  <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
  <220>
  <223> At position 10, Xaa is an azetidine residue
<400> 914
  Phe Glu Trp Thr Pro Gly Trp Tyr Gln Xaa Tyr Ala Leu Pro Leu
                               10
  1 5
  <210> 915
  <211> 15
  <212> PRT
  <213> Artificial Sequence
  <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
  <400> 915
  Phe Glu Trp Thr Pro Gly Tyr Tyr Gln Pro Tyr Ala Leu Pro Leu
                            10
                  5
  <210> 916
  <211> 15
   <212> PRT
   <213> Artificial Sequence
   <223> Description of Artificial Sequence: IL-1 ANTAGONIST
```

PEPTIDE

```
<220>
<223> At position 10, Xaa is an azetidine residue
<400> 916
Phe Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Tyr Ala Leu Pro Leu
                  5
<210> 917
<211> 21
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 1, Xaa is A, D, E, F, G, K, Q, S, T, V
      or Y
<220>
<223> At position 2, Xaa is A, D, G, I, N, P, S, T, V or
<220>
<223> At position 3, Xaa is A, D, G, L, N, P, S, T, W or
      Y
<223> At position 4, Xaa is A, D, E, F, L, N, R, V or Y
<223> At position 5, Xaa is A, D, E, Q, R, S or T
<223> At position 6, Xaa is H, I, L, P, S, T or W
<220>
<223> At position 7, Xaa is A, E, F, K, N, Q, R, S or Y
<220>
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<223> At position 8, Xaa is D, E, F, Q, R, T or W <220>

<223> At position 9, Xaa is A, D, P, S, T or W

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<220>
<223> At position 10, Xaa is A, D, G, K, N, Q, S or T
<220>
<223> At position 11, Xaa is A, E, L, P, S, T, V or Y
<220>
<223> At position 12, Xaa is V, L, I, E, P, G, Y, M, T
     or D
<220>
<223> At position 13, Xaa is Y, W or F
<220>
<223> At position 14, Xaa is E, F, V, W or Y
<220>
<223> At position 16, Xaa is P or an azetidine residue
<223> At position 18, Xaa is S, A, V or L
<223> At position 19, Kaa is M, F, V, R, Q, K, T, S, D,
     L, I or E
<220>
<223> At position 20, Xaa is Q or P
<400> 917
15
                                  10
Tyr Xaa Xaa Xaa Leu
            20
 <210> 918
 <211> 21
 <212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
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PCT/US99/25044

WO 00/24782 <400> 918 Thr Ala Asn Val Ser Ser Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro 10 Tyr Ala Leu Pro Leu 20 <210> 919 <211> 18 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE <400> 919 Ser Trp Thr Asp Tyr Gly Tyr Trp Gln Pro Tyr Ala Leu Pro Ile Ser 10 5 . 1 Gly Leu

<210> 920 <211> 21 <212> PRT <213> Artificial Sequence

<220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 920 Glu Thr Pro Phe Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro 15 5

Tyr Ala Leu Pro Leu 20

<210> 921 <211> 21 <212> PRT

<213> Artificial Sequence <220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE <400> 921 Glu Asn Thr Tyr Ser Pro Asn Trp Ala Asp Ser Met Tyr Trp Gln Pro 10 Tyr Ala Leu Pro Leu 20 <210> 922 <211> 21 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE <400> 922 Ser Val Gly Glu Asp His Asn Phe Trp Thr Ser Glu Tyr Trp Gln Pro 10 1 5 Tyr Ala Leu Pro Leu 20 <210> 923 <211> 21 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE <400> 923 Asp Gly Tyr Asp Arg Trp Arg Gln Ser Gly Glu Arg Tyr Trp Gln Pro 10 5 1

Tyr Ala Leu Pro Leu 20

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<210> 924
<211> 15
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST
<400> 924
Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro Tyr Ala Leu Pro Leu
                                    10
                5
<210> 925
<211> 13
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 925
Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro Tyr Asn His
                                    10
                  5
<210> 926
<211> 13
<212> PRT
<213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <220>
 <223> At position 10, Xaa is an azetidine residue
 <400> 926 ···
 Phe Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr Asn His
                                      10
```

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<210> 927
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST
      PEPTIDE
<400> 927
Glu Trp Thr Pro Gly Tyr Trp Gln Pro Tyr Asn His
                  5
<210> 928
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 10, Xaa is an azetidine residue
<400> 928
Phe Glu Trp Thr Pro Gly Trp Tyr Gln Xaa Tyr
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<210> 929
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <220>
<223> At position 10, Xaa is an azetidine residue
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<400> 929
Ala Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr
                  5
<210> 930
<211> 11
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <220>
 <223> At position 10, Xaa is an azetidine residue
 <400> 930
 Phe Ala Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr
  ຸ 1
<210> 931
 <211> 11
 <212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <220>
 <223> At position 10, Xaa is an azetidine residue
 <400> 931
  Phe Glu Ala Thr Pro Gly Tyr Trp Gln Xaa Tyr
                                      10
                    5
  <210> 932
  <211> 11
  <212> PRT
  <213> Artificial Sequence
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<220>

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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<220>
<223> At position 10, Xaa is an azetidine residue
<400> 932
Phe Glu Trp Ala Pro Gly Tyr Trp Gln Xaa Tyr
               5
<210> 933
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
    PEPTIDE
<220>
<223> At position 10, Xaa is an azetidine residue
<400> 933
Phe Glu Trp Thr Ala Gly Tyr Trp Gln Xaa Tyr
                5 . 10
<210> 934
<211> 11
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <220>
 <223> At position 10, Xaa is an azetidine residue
 Phe Glu Trp Thr Pro Ala Tyr Trp Gln Xaa Tyr
  1 ... 5
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<210> 935
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 10, Xaa is an azetidine residue
<400> 935
Phe Glu Trp Thr Pro Gly Ala Trp Gln Xaa Tyr
                  5
<210> 936
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 10, Xaa is an azetidine residue
<400> 936
Phe Glu Trp Thr Pro Gly Tyr Ala Gln Xaa Tyr
<210> 937
<211> 11
<212> PRT
<213> Artificial Sequence
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <220>
```

<223> At position 10, Xaa is an azetidine residue

```
<400> 937
Phe Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Ala
1 5 10
```

5

<210> 939
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE
<220>
<223> At position 5, D amino acid residue
<220>
<223> At position 10, Xaa is an azetidine residue

Phe Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr

<210> 940 <211> 10 <212> PRT

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<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<220>
<223> At position 10, Xaa is an azetidine residue
<400> 940
Phe Glu Trp Thr Gly Tyr Trp Gln Xaa Tyr
                 5
<210> 941
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<220>
<223> At position 5, Xaa is a pipecolic acid residue
<220>
<223> At position 10, Xaa is an azetidine residue
<400> 941
Phe Glu Trp Thr Xaa Gly Tyr Trp Gln Xaa Tyr
<210> 942
<211> 11
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
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<223> At position 6, Xaa is an aminoisobutyric acid

residue

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<220>
 <223> At position 10, Xaa is an azetidine residue
 <400> 942
 Phe Glu Trp Thr Pro Xaa Tyr Trp Gln Xaa Tyr
                   5
 <210> 943
 <211> 11
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <220>
 <223> At position 6, Kaa is a sarcosine residue
 <223> At position 10, Xaa is an azetidine residue
 <400> 943
 Phe Glu Trp Thr Pro Xaa Trp Tyr Gln Xaa Tyr
                    5
  <210> 944
  <211> 11
  <212> PRT
<213> Artificial Sequence
  <220>
  <223> Description of Artificial Sequence:IL-1 ANTAGONIST
        PEPTIDE
  <220>
  <223> At position 5, Xaa is a sarcosine residue
  <220>
  <223> At position 10, Kaa is an azetidine residue
  <400> 944
  Phe Glu Trp Thr Xaa Gly Tyr Trp Gln Xaa Tyr
```

1 5 10

<210> 945

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<220>

<223> At position 10, Xaa is an azetidine residue

<400> 945

Phe Glu Trp Thr Pro Asn Tyr Trp Gln Xaa Tyr
1 5 10

<210> 946

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<220>

<223> At position 5, D amino acid residue

<220>

<223> At position 10, Xaa is an azetidine residue

<400> 946

Phe Glu Trp Thr Pro Val Tyr Trp Gln Xaa Tyr 1 5 10

<210> 947

<211> 11 ...

<212> PRT

<213> Artificial Sequence

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<220>
  <223> Description of Artificial Sequence: IL-1 ANTAGONIST
        PEPTIDE
  <220>
  <223> At position 10, Xaa is an azetidine residue
  <400> 947
  Phe Glu Trp Thr Val Pro Tyr Trp Gln Xaa Tyr
                   5
  <210> 948
  <211> 11
  <212> PRT
  <213> Artificial Sequence
 <220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
        PEPTIDE
  <220>
  <223> At position 1, Xaa is acetylated phe
  <220>
  <223> At position 10, Xaa is an azetidine residue
  ≼400> 948
  Phe Glu Trp Thr Pro Gly Trp Tyr Gln Xaa Tyr
                   5
  <210> 949
  <211> 11
  <212> PRT
  <213> Artificial Sequence
   <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
   <220>
   <223> At position 1, Xaa is acetylated phe
   <220>
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<223> At position 10, Xaa is an azetidine residue

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<400> 949
Phe Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr
       5
<210> 950
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<220>
<223> At position 1, Xaa=1-naphthylalanine
<223> At position 10, Xaa is an azetidine residue
<400> 950
Xaa Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Tyr
            5
<210> 951
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<220>
<223> At position 10, Xaa is an azetidine residue
<400> 951
Tyr Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Tyr
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<210> 952 <211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<220>

<223> At position 10, Xaa is an azetidine residue

<400> 952

Phe Glu Trp Val Pro Gly Tyr Tyr Gln Xaa Tyr
1 5 10

<210> 953

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<220>

<223> At position 10, Xaa is an azetidine residue

<400> 953

Phe Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Tyr
1 5 10

<210> 954

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<220>

<223> At position 10, Xaa is an azetidine residue

<400> 954

Phe Glu Trp Thr Pro Ser Tyr Tyr Gln Xaa Tyr

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<210> 955 <211> 11 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE <220> <223> At position 10, Xaa is an azetidine residue <400> 955 Phe Glu Trp Thr Pro Asn Tyr Tyr Gln Xaa Tyr 5 <210> 956 <211> 12 <212> PRT <213> Artificial Sequence <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE <220> <223> At position 5, Xaa=naphthylalanine <400> 956 Ser His Leu Tyr Xaa Gln Pro Tyr Ser Val Gln Met 5

<210> 957 <211> 12 <212> PRT <213> Artificial Sequence

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<220>
<223> At position 5, Xaa=naphthylalanine
<400> 957
Thr Leu Val Tyr Xaa Gln Pro Tyr Ser Leu Gln Thr
                  5
<210> 958
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 5, Xaa=naphthylalanine
<400> 958
Arg Gly Asp Tyr Xaa Gln Pro Tyr Ser Val Gln Ser
<210> 959
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 5, Xaa=naphthylalanine
<400> 959
Asn Met Val Tyr Xaa Gln Pro Tyr Ser Ile Gln Thr
                  5
  1
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<210> 960 <211> 9

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<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 960
Val Tyr Trp Gln Pro Tyr Ser Val Gln
                 5
<210> 961
<211> 9
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
<220>
<223> At position 3, Xaa=naphthylalanine
<400> 961
Val Tyr Xaa Gln Pro Tyr Ser Val Gln
                  5
<210> 962
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
<220>
<223> At position 7, Xaa is an azetidine residue
<400> 962
 Thr Phe Val Tyr Trp Gln Xaa Tyr Ala Leu Pro Leu
                   5
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<210> 963
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<220>
<223> At position 10, Xaa is an azetidine residue
<220>
<223> At position 11, Xaa =p-benzoyl-L-phenylalanine
<400> 963
Phe Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Xaa
                  5
<210> 964
<211> 11
<212> PRT
<213> Artificial Sequence
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <220>
 <223> At position 1, Xaa=acetylated phe
 <223> At position 10, Xaa is an azetidine residue
 <223> At position 11, Xaa=p-benzoyl-L-phenylalanine
 <400> 964
 Xaa Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Xaa
                   5
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<210> 965 <211> 11

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<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 8, Xaa=p-benzoyl-L-phenylalanine
<220>
<223> At position 10, Xaa is an azetidine residue
<400> 965
Phe Glu Trp Thr Pro Gly Tyr Xaa Gln Xaa Tyr
                                     10
                  5
<210> 966
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 1, Xaa=acetylated phe
<223> At position 8, Xaa=p-benzoyl-L-phenylalanine
<220>
<223> At position 10, Xaa is an azetidine residue
 <400> 966
 Phe Glu Trp Thr Pro Gly Tyr Xaa Gln Xaa Tyr
                   5
 <210> 967
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<210> 967 <211> 11 <212> PRT ---<213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 7, Xaa=p-benzoyl-L-phenylalanine
<220>
<223> At position 10, Xaa is an azetidine residue
<400> 967
Phe Glu Trp Thr Pro Gly Xaa Tyr Gln Xaa Tyr
                 5
<210> 968
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
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PEPTIDE

<220> <223> At position 1, Xaa=acetylated phe

<223> At position 7, Xaa=p-benzoyl-L-phenylalanine

<223> At position 10, Xaa is an azetidine residue <400> 968 Phe Glu Trp Thr Pro Gly Xaa Tyr Gln Xaa Tyr

5

<210> 969 <211> 11 <212> PRT <213> Artificial Sequence <220>

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

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<220>
<223> At position 1, Xaa=acetylated phe
<223> At position 3, Xaa=p-benzoyl-L-phenylalanine
<220>
<223> At position 10, Xaa is an azetidine residue
<400> 969
Phe Glu Xaa Thr Pro Gly Tyr Tyr Gln Xaa Tyr
<210> 970
<211> 11
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 1, Xaa=acetylated phe
<220>
<223> At position 3, Xaa=p-benzoyl-L-phenylalanine
<223> At position 10, Xaa is an azetidine residue
Phe Glu Xaa Thr Pro Gly Tyr Tyr Gln Xaa Tyr
                  5
<210> 971
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
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<220>
<223> At position 1, Xaa=p-benzoyl-L-phenylalanine
<223> At position 10, Xaa is an azetidine residue
<400> 971
Xaa Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Tyr
<210> 972
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 1, Xaa=acetylated
      p-benzoyl-L-phenylalanine
<223> At position 10, Xaa is an azetidine residue
<400> 972
Xaa Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Tyr
                  5
<210> 973
<211> 9
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 973
Val Tyr Trp Gln Pro Tyr Ser Val Gln
         , 5
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<210> 974
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<400> 974
Arg Leu Val Tyr Trp Gln Pro Tyr Ser Val Gln Arg
       . 5
<210> 975
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<223> At position 5, Xaa=naphthylalanine
<400> 975
Arg Leu Val Tyr Xaa Gln Pro Tyr Ser Val Gln Arg
                                   10
                 5
<210> 976
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 976
Arg Leu Asp Tyr Trp Gln Pro Tyr Ser Val Gln Arg
                  5
  1
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<210> 977
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 977
Arg Leu Val Trp Phe Gln Pro Tyr Ser Val Gln Arg
                  5
                                     10
<210> 978
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 978
Arg Leu Val Tyr Trp Gln Pro Tyr Ser Ile Gln Arg
                  5
  1
<210> 979
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<223> At position 1, Xaa=D or Y
<223> At position 3, Xaa=D or S
 <220>
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<223> At position 4, Xaa=S, T or A
<220>
<223> At position 5, Xaa=S or W
<220>
<223> At position 6, Xaa=S or Y
<220>
<223> At position 7, Xaa=D, Q, E or V
<223> At position 8, Xaa=N, S, K, H or W
<220>
<223> At position 9, Xaa=F or L
<220>
<223> At position 10, Xaa=D, N, S or L
<220>
<223> At position 11, Xaa=L, I, Q, M or A
<400> 979
Xaa Asn Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa
                                     10
                  5
  1
<210> 980
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 980
Asp Asn Ser Ser Trp Tyr Asp Ser Phe Leu Leu
  1
                  5
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<210> 981 <211> 11 ... <212> PRT <213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<400> 981
Asp Asn Thr Ala Trp Tyr Glu Ser Phe Leu Ala
1 5
<210> 982
<211> 11
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<400> 982
Asp Asn Thr Ala Trp Tyr Glu Asn Phe Leu Leu
     5
<210> 983
<211> 17
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<400> 983
Pro Ala Arg Glu Asp Asn Thr Ala Trp Tyr Asp Ser Phe Leu Ile Trp
                                    10
                  5
Сув
<210> 984
<211> 17 ...
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<212> PRT

<213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 984
Thr Ser Glu Tyr Asp Asn Thr Thr Trp Tyr Glu Lys Phe Leu Ala Ser
                5
                                   10
Gln
<210> 985
<211> 17
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 985
Ser Gln Ile Pro Asp Asn Thr Ala Trp Tyr Gln Ser Phe Leu Leu His
                                                        15
                  5
                                    10
Gly
<210> 986
<211> 17
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <400> 986
Ser Pro Phe Ile Asp Asn Thr Ala Trp Tyr Glu Asn Phe Leu Leu Thr
                                                         15
                                     10
                   5
  1
```

Tyr

```
<210> 987
 <211> 17
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:IL-1 ANTAGONIST
       PEPTIDE
 <400> 987
 Glu Gln Ile Tyr Asp Asn Thr Ala Trp Tyr Asp His Phe Leu Leu Ser
                 5
                                     10
 Tyr
 <210> 988
 <211> 17
 <212> PRT
  <213> Artificial Sequence
 <220>
  <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
  <400> 988
Thr Pro Phe Ile Asp Asn Thr Ala Trp Tyr Glu Asn Phe Leu Leu Thr
                                     10
                    5
  Tyr
  <210> 989
  <211> 17
  <212> PRT
  <213> Artificial Sequence
· <220>
  <223> Description of Artificial Sequence:IL-1 ANTAGONIST
        PEPTIDE
  <400> 989
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Thr Tyr Thr Tyr Asp Asn Thr Ala Trp Tyr Glu Arg Phe Leu Met Ser

1 5 10 15

Tyr

<210> 990

<211> 17

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<400> 990

Thr Met Thr Gln Asp Asn Thr Ala Trp Tyr Glu Asn Phe Leu Leu Ser

1 5 10 15

Tyr

<210> 991

<211> 17

<212> PRT

<213> Artificial Sequence

<220>

<400> 991

Thr Ile Asp Asn Thr Ala Trp Tyr Ala Asn Leu Val Gln Thr Tyr Pro 1 5 10 15

Gln

<210> 992

<211> 17 ...

<212> PRT

<213> Artificial Sequence

<220>

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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 992
Thr Ile Asp Asn Thr Ala Trp Tyr Glu Arg Phe Leu Ala Gln Tyr Pro
                  5
                                     10
Asp
<210> 993
<211> 17
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 993
His Ile Asp Asn Thr Ala Trp Tyr Glu Asn Phe Leu Leu Thr Tyr Thr
                                     10
                  5
Pro
<210> 994
<211> 17
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST
      PEPTIDE
<400> 994
Ser Gln Asp Asn Thr Ala Trp Tyr Glu Asn Phe Leu Leu Ser Tyr Lys
                                                          15
                                     10
                  5
Ala
```

```
<210> 995
 <211> 17
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:IL-1 ANTAGONIST
       PEPTIDE
 <400> 995
 Gin Ile Asp Asn Thr Ala Trp Tyr Glu Arg Phe Leu Leu Gln Tyr Asn
                                 10
                  5
 Ala
 <210> 996
 <211> 17
 <212> PRT
 <213> Artificial Sequence
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 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <400> 996
Asn Gln Asp Asn Thr Ala Trp Tyr Glu Ser Phe Leu Leu Gln Tyr Asn
                                     10
                  5
 Thr
  <210> 997
  <211> 17
  <212> PRT
  <213> Artificial Sequence
  <220>
  <223> Description of Artificial Sequence: IL-1 ANTAGONIST
        PEPTIDE
  <400> 997
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WO 00/24782
Thr Ile Asp Asn Thr Ala Trp Tyr Glu Asn Phe Leu Leu Asn His Asn
                  5
                                     10
Leu
<210> 998
<211> 17
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL·1 ANTAGONIST
      PEPTIDE
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<400> 998

His Tyr Asp Asn Thr Ala Trp Tyr Glu Arg Phe Leu Gln Gln Gly Trp 5 10

His

<210> 999 <211> 21 <212> PRT <213> Artificial Sequence <223> Description of Artificial Sequence: IL-1 ANTAGONIST PEPTIDE

<400> 999 Glu Thr Pro Phe Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro 15 10

Tyr Ala Leu Pro Leu 20

<210> 1000 <211> 21 ... <212> PRT . <213> Artificial Sequence

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<220>
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      PEPTIDE
<400> 1000
Tyr Ile Pro Phe Thr Trp Glu Glu Ser Asn Ala Tyr Tyr Trp Gln Pro
                                    10
Tyr Ala Leu Pro Leu
           20
<210> 1001
<211> 21
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <400> 1001
Asp Gly Tyr Asp Arg Trp Arg Gln Ser Gly Glu Arg Tyr Trp Gln Pro
                   5
 1
 Tyr Ala Leu Pro Leu
              20
 <210> 1002
 <211> 10
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <223> At position 1, Xaa=phosphotyrosine
 <220>
  <223> At position 2, Xaa=naphthylalanine
```

<220>

```
<223> At position 3, Xaa=phosphotyrosine
<223> At position 5, Xaa is an azetidine residue
<400> 1002
Xaa Xaa Xaa Gln Xaa Tyr Ala Leu Pro Leu
                 5
<210> 1003
<211> 21
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 1003
Thr Ala Asn Val Ser Ser Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro
                                     10
                  5
Tyr Ala Leu Pro Leu
             20
<210> 1004
<211> 15
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 10, Xaa=azetidine
<400> 1004
Phe Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr Ala Leu Pro Leu
                                      10
                  5
```

<210> 1005

```
<211> 19
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<400> 1005
Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro Tyr Ala Leu Pro Leu Ser
                                  10
            5
Asp Asn His
<210> 1006
<211> 15
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 10, Xaa=azetidine
<400> 1006
Phe Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Tyr Ala Leu Pro Leu
                                    10
                5
<210> 1007
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
 <220>
 <223> At position 10, Xaa=azetidine
```

<400> 1007

```
<210> 1008
<211> 11
<212> PRT
<213> Artificial Sequence
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
     PEPTIDE
<220>
<223> At position 1, Xaa=acetylated phe
<223> At position 10, Xaa=azetidine
<400> 1008
Phe Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr
                 5
<210> 1009
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 1, Xaa=acetylated phe
<220>
<223> At position 10, Xaa=azetidine
<400> 1009
Phe Glu Trp Thr Pro Gly Trp Tyr Gln Xaa Tyr
```

Phe Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr

```
<211> 11
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 1, Xaa=acetylated phe
<223> At position 10, Xaa=azetidine
 <400> 1010
 Phe Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Tyr
                   5
 <210> 1011
 <211> 11
 <212> PRT
 <213> Artificial Sequence
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 <223> Description of Artificial Sequence: IL-1 ANTAGONIST
       PEPTIDE
 <220>
<223> At position 1, Xaa=acetylated phe
 <220>
 <223> At position 10, Xaa=azetidine
 <400> 1011
 Phe Glu Trp Thr Pro Ala Tyr Trp Gln Xaa Tyr
                    5
  <210> 1012
  <211> 11
  <212> PRT
  <213> Artificial Sequence
  <220>
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST

PEPTIDE

<220>
<223> At position 1, Xaa=acetylated phe

<220>
<223> At position 10, Xaa=azetidine

<400> 1012
Phe Glu Trp Thr Pro Ala Trp Tyr Gln Xaa Tyr
1 5 10

<210> 1013

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:IL-1 ANTAGONIST PEPTIDE

<220>

<223> At position 1, Xaa=acetylated phe

<220>

<223> At position 10, Xaa=azetidine

<400> 1013

Phe Glu Trp Thr Pro Ala Tyr Tyr Gln Xaa Tyr
1 5 10

<210> 1014

<211> 15

<212> PRT

<213> Artificial Sequence

<220>

<220>

<223> At position 10, Xaa=azetidine

<400> 1014

Phe Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Tyr Ala Leu Pro Leu
1 5 10 15

<210> 1015

<211> 15

<212> PRT

<213> Artificial Sequence

<220>

<220>

<223> At position 10, Xaa=azetidine

<400> 1015

Phe Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr Ala Leu Pro Leu
1 5 10 15

<210> 1016

<211> 15

<212> PRT

<213> Artificial Sequence

<220>

<220>

<223> At position 10, Xaa=azetidine

<400> 1016

Phe Glu Trp Thr Pro Gly Trp Tyr Gln Xaa Tyr Ala Leu Pro Leu
1 5 10 15

<210> 1017

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST

PEPTIDE

<400> 1017

Thr Ala Asn Val Ser Ser Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro 1 5 10 15

Tyr Ala Leu Pro Leu 20

<210> 1018

<211> 11

<212> PRT

<213> Artificial Sequence

<220>

<220>

<223> At position 1, Xaa=acetylated phe

<220>

<223> At position 10, Xaa=azetidine

<400> 1018

Phe Glu Trp Thr Pro Gly Tyr Trp Gln Xaa Tyr
1 5 10

<210> 1019

<211> 11

<212> PRT

<213> Artificial Sequence

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<220>

<223> At position 1, Xaa=acetylated phe

<220>

<223> At position 10, Xaa=azetidine

<400> 1019

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Phe Glu Trp Thr Pro Gly Trp Tyr Gln Xaa Tyr
1 5 10
```

<210> 1021 <211> 11 <212> PRT <213> Artificial Sequence

<220>
<223> At position 1, Xaa=acetylated phe
<220>

Phe Glu Trp Thr Pro Gly Tyr Tyr Gln Xaa Tyr

5

<223> At position 6, D amino acid residue
.
<220>

<223> At position 10, Xaa=azetidine

<400> 1021
Phe Glu Trp Thr Pro Ala Tyr Trp.Gln Xaa Tyr
1 5 10

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<210> 1022
<211> 11
<212> PRT
<213> Artificial Sequence
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PEPTIDE
<220>
<223> At position 1, Xaa=acetylated phe
<223> At position 6, D amino acid residue
<220>
<223> At position 10, Xaa=azetidine
<400> 1022
Phe Glu Trp Thr Pro Ala Trp Tyr Gln Xaa Tyr
                  5
<210> 1023
<211> 11
<212> PRT
<213> Artificial Sequence
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<223> Description of Artificial Sequence: IL-1 ANTAGONIST
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<220>
<223> At position 1, Xaa=acetylated phe
<220>
<223> At position 6, D amino acid residue
<220>
 <223> At position 10, Xaa=azetidine
 <400> 1023
 Phe Glu Trp Thr Pro Ala Tyr Tyr Gln Xaa Tyr
                  5
   1
```

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<210> 1024
<211> 20
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EPO MIMETIC
     PEPTIDE
<400> 1024
Gly Gly Leu Tyr Leu Cys Arg Phe Gly Pro Val Thr Trp Asp Cys Gly
                5
Tyr Lys Gly Gly
             20
<210> 1025
<211> 20
<212> PRT
<213> Artificial Sequence
<220>
 <223> Description of Artificial Sequence: EPO MIMETIC
     PEPTIDE
 <400> 1025
 Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys
                                    10
                 5
  1
 Pro Gln Gly Gly
              20
 <210> 1026
 <211> 20
 <212> PRT
 <213> Artificial Sequence
<220>
 <223> Description of Artificial Sequence: EPO-MIMETIC
       PEPTIDE
```

<400> 1026

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Gly Gly Asp Tyr His Cys Arg Met Gly Pro Leu Thr Trp Val Cys Lys
1 5 10 15
```

Pro Leu Gly Gly 20

<210> 1027

<211> 13

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EPO MIMETIC PEPTIDE

<400> 1027

Cys Gly Arg Glu Cys Pro Arg Leu Cys Gln Ser Ser Cys

1 5 10

<210> 1028

<211> 13

<212> PRT

<213> Artificial Sequence

<220>

<400> 1028

Cys Asn Gly Arg Cys Val Ser Gly Cys Ala Gly Arg Cys
1 5 10

<210> 1029

<211> 20

<212> PRT

<213> Artificial Sequence

<220>

<400> 1029

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Val Gly Asn Tyr Met Cys His Phe Gly Pro Ile Thr Trp Val Cys Arg
                                     10
Pro Gly Gly Gly
<210> 1030
<211> 20
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: EPO MIMETIC
      PEPTIDE
<400> 1030
Gly Gly Val Tyr Ala Cys Arg Met Gly Pro Ile Thr Trp Val Cys Ser
                                      10
Pro Leu Gly Gly
              20
 <210> 1031
 <211> 5
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: VEGF ANTAGONIST
       PEPTIDE
 <400> 1031
 Cys Asn Gly Arg Cys
   1
  <210> 1032
  <211> 9
  <212> PRT
  <213> Artificial Sequence
  <220>
  <223> Description of Artificial Sequence: TPO MIMETIC
```

```
<400> 1032
Cys Asp Cys Arg Gly Asp Cys Phe Cys
      5
<210> 1033
<211> 20
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: EPO MIMETIC
 <400> 1033
 Ile Glu Gly Pro Thr Leu Arg Gln Trp Leu Ala Ala Arg Ala Gly Gly
                                    10
             5
 Gly Gly Gly Phe
 <210> 1034
 <211> 26
 <212> PRT
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<223> Description of Artificial Sequence: EPO MIMETIC
 <400> 1034
 Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys
 Pro Gln Gly Gly Gly Gly Gly Phe
              20
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<210> 1035 <211> 19 <212> PRT <213> Artificial Sequence

<220>
<223> Description of Artificial Sequence: EPO MIMETIC

<400> 1035

Val Gly Asn Tyr Met Ala His Met Gly Pro Ile Thr Trp Val Cys Arg

1 5 10 15

Pro Gly Gly

<210> 1036

<211> 18

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EPO MIMETIC

<400> 1036

Gly Gly Thr Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys
1 5 10 15

Pro Gln

<210> 1037

<211> 20

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EPO MIMETIC

**<400> 1037** 

Gly Gly Leu Tyr Ala Cys His Met Gly Pro Met Thr Trp Val Cys Gln
1 5 10 15

Pro Leu Arg Gly

20

<210> 1038

<211> 22 ...

<212> PRT

<213> Artificial Sequence

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<220>
<223> Description of Artificial Sequence: EPO MIMETIC
<400> 1038
Thr Ile Ala Gln Tyr Ile Cys Tyr Met Gly Pro Glu Thr Trp Glu Cys
                                   10
                 5
Arg Pro Ser Pro Lys Ala
            20
<210> 1039 ·
<211> 13
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<223> Description of Artificial Sequence: EPO MIMETIC
<400> 1039
Tyr Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys
                                    10
 1
<210> 1040
<211> 11
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: EPO MIMETIC
      PEPTIDE
<400> 1040
Tyr Cys His Phe Gly Pro Leu Thr Trp Val Cys
                 5
 1
 <210> 1041
<211> 12
 <212> PRT
 <213> Artificial Sequence
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<220>

<223> Description of Artificial Sequence: EPO MIMETIC PEPTIDE

<400> 1041 Ser Cys His Phe Gly Pro Leu Thr Trp Val Cys Lys 5

<210> 1042

<211> 40

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: EPO MIMETIC PEPTIDE

<400> 1042

10 5 ·

Pro Xaa Xaa Xaa Xaa Xaa Xaa Thr Trp Xaa Xaa Xaa Xaa Xaa Xaa 25 20

Xaa Xaa Xaa Xaa Xaa Xaa Xaa 35

<210> 1043

<211> 5

<212> PRT

<213> Artificial Sequence

<223> Description of Artificial Sequence: EPO MIMETIC PEPTIDE

<400> 1043

Asp Leu Xaa Xaa Leu 1

<210> 1044

<211> 12

<212> PRT

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<213> Artificial Sequence
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<220>

<223> Description of Artificial Sequence: INTEGRIN BINDING PEPTIDE

<400> 1044

Arg Thr Asp Leu Asp Ser Leu Arg Thr Tyr Thr Leu
1 5 10

<210> 1045

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TNF ANTAGONIST

<400> 1045

Phe Gly Gly Gly Gly Asp Phe Leu Pro His Tyr Lys Asn Thr Ser 1 5 10 15

Leu Gly His Arg Pro 20

<210> 1046

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: TNF ANTAGONIST

<400> 1046

Asp Phe Leu Pro His Tyr Lys Asn Thr Ser Leu Gly His Arg Pro Gly
1 5 10 15

Gly Gly Gly Phe 20

<210> 1047

<211> 21

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<212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: IL-1 ANTAGONIST <400> 1047

Phe Gly Gly Gly Gly Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro . 5

Tyr Ala Leu Pro Leu 20

<210> 1048 <211> 21 <212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST

<400> 1048

Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro Tyr Ala Leu Pro Leu Gly 10 5

Gly Gly Gly Phe 20

<210> 1049 <211> 25 <212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: VEGF ANTAGONIST

<400> 1049

Phe Gly Gly Gly Gly Val Glu Pro Asn Cys Asp Ile His Val Met 10 5

Trp Glu Trp Glu Cys Phe Glu Arg Leu ... 20

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<210> 1050
<211> 25
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: VEGF ANTAGONIST
<400> 1050
Val Glu Pro Asn Cys Asp Ile His Val Met Trp Glu Trp Glu Cys Phe
                                  10
                5
Glu Arg Leu Gly Gly Gly Gly Phe
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<210> 1051
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<212> PRT
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Phe Gly Gly Gly Gly Cys Thr Thr His Trp Gly Phe Thr Leu Cys
                 5
<210> 1052
<211> 16
 <212> PRT
 <213> Artificial Sequence
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 <400> 1052
 Cys Thr Thr His Trp Gly Phe Thr Leu Cys Gly Gly Gly Gly Phe
                                   10
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<210> 1053 <211> 10

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<212> PRT
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<223> Description of Artificial Sequence: INTEGRIN
     BINDING PEPTIDE
<400> 1053
Arg Thr Asp Leu Asp Ser Leu Arg Thr Tyr
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<210> 1054
<211> 9
<212> PRT
<213> Artificial Sequence
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<223> Description of Artificial Sequence: INTEGRIN
      BINDING PEPTIDE
<400> 1054
Arg Thr Asp Leu Asp Ser Leu Arg Thr
                 5
<210> 1055
<211> 757
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<213> Artificial Sequence
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     Met Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu Leu
                                         10
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ctg ggg gga ccg tca gtc ttc ctc ttc ccc cca aaa ccc aag gac acc Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr

5

20 25 30

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	-								-					Val			
				35	•				40					45	-		
a	gc	cac	gaa	gac	cct	gag	gtc	aag	ttc	aac	tgg	tac	gtg	gac	ggc	gtg	192
	-		-	-										Asp			
			50					55					60				
																-	
g	ag.	gtg	cat	aat	gcc	aag	aca	aag	ccg	cgg	gag	gag	cag	tac	aac	agc	240
G	lu	Val	His	Asn	Ala	Lys	Thr	Lys	Pro	Arg	Glu	Glu	Gln	Tyr	Asn	Ser	
		65					70					75					
														gac			288
T	hr	Tyr	Arg	Val	Val	Ser	Val	Leu	Thr	Val	Leu	His	Gln	Asp	Trp	Leu	
	80					85					90					95	
																•	
														ctc			336
A	sn	Gly	Lys	Glu	Tyr	Lys	Cys	Lys	Val		Asn	Lys	Ala	Leu		Ala	
					100					105					110		
•	•																204
														cga			384
P	,LO	Ile	Glu	Lys	Thr	Ile	Ser	Lys		Lys	Gly	Gln	Pro	Arg	Glu	Pro	
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																	432
														aag			472
G	ln	Val	_	Thr	Leu	Pro	Pro		Arg	ASD	GIU	Leu		Lys	ABII	GIII	
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													200	aac	ato	acc	480
g	rtc	agc	ctg	acc	tgc	ctg	gtc	aaa	ggc	Dha	Tur	Pro	Ser	gac	Tle	Ala	
V	aı		Leu	Thr	Сув	ren	150	nys	GIY	FIIC	1 Y L	155	501	Asp			
		145					150					100					
_		~n~	+ ~ ~	a=a	200	22 t	aaa	сал	cca	gag	aac	aac	tac	aag	acc	acg	528
7	7-1	Glu	Tra	Glu	Ser	lan	'G1v	Gln	Pro	Glu	Asn	Asn	Tyr	Lys	Thr	Thr	
	.60	GIU	ענונ	GIU	Der	165	013	<b>V</b>	•••		170		-	-		175	
-	.00																
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ב	220	Pro	Val	Leu	Asp	Ser	Asp	Glv	Ser	Phe	Phe	Leu	Tyr	Ser	Lys	Leu	
•		FLO	141	200	180					185					190		
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η	 Thr	Val	Asn	Lvs	Ser	Ara	Trp	Gln	Gln	Gly	Asn	Val	Phe	Ser	Cys	Ser	
•			E	195		3			200	-				205	ı		
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7	Val	Met	His	Glu	Ala	Leu	His	Asn	His	Tyr	Thr	Gln	Lys	Ser	Leu	Ser	

210 215 220

ctg tct ccg ggt aaa ggt gga ggt ggt ggt gac ttc ctg ccg cac tac 720
Leu Ser Pro Gly Lys Gly Gly Gly Gly Asp Phe Leu Pro His Tyr
225 230 235

aaa aac acc tct ctg ggt cac cgt ccg taatggatcc 757
Lys Asn Thr Ser Leu Gly His Arg Pro
240 245

<210> 1056

<211> 248

<212> PRT

<213> Artificial Sequence

<400> 1056

Met Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu Leu Leu

1 5 10 15

Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr Leu 20 25 30

Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp Val Ser

His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly Val Glu 50 55 60

Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Glu Tyr Asn Ser Thr
65 70 75 80

Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln Asp Trp Leu Asn 85 90 95

Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu Pro Ala Pro 100 105 110

Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro Gln 115 120 125

Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn Gln Val 130 135 140

Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala Val 145 150 155 160

Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr Thr Pro 165 170 175

Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu Thr 180 185 190

Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser Val 195 200 205

Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser Leu 210 220

Ser Pro Gly Lys Gly Gly Gly Gly Gly Asp Phe Leu Pro His Tyr Lys 225 230 235 240

Asn Thr Ser Leu Gly His Arg Pro 245

<210> 1057

<211> 761

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:TNF-ALPH INHIBITOR Fc

<220>

<221> CDS

<222> (4)..(747)

<400> 1057

cat atg gac ttc ctg ccg cac tac aaa aac acc tct ctg ggt cac cgt 48

Met Asp Phe Leu Pro His Tyr Lys Asn Thr Ser Leu Gly His Arg

1 5 10 15

ccg ggt gga ggc ggt ggg gac aaa act cac aca tgt cca cct tgc cca 96
Pro Gly Gly Gly Gly Asp Lys Thr His Thr Cys Pro Pro Cys Pro
20 25 30

gca cct gaa ctc ctg ggg gga ccg tca gtt ttc ctc ttc ccc cca aaa 144
Ala Pro Glu Leu Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys
... 35 40 45

ccc aag gac acc ctc atg atc tcc cgg acc cct gag gtc aca tgc gtg 192

Pro	Lys	Asp 50	Thr	Leu	Met	Ile	Ser 55	Arg	Thr	Pro	Glu	Val 60	Thr	Сув	Val	
					cac His											240
					gtg Val 85											288
cag Gln	tac Tyr	aac Asn	agc Ser	acg Thr 100	tac Tyr	cgt Arg	gtg Val	gtc Val	agc Ser 105	gtc Val	ctc Leu	acc Thr	gtc Val	ctg Leu 110	cac His	336
cag Gln	gac Asp	tgg Trp	ctg Leu 115	aat Asn	ggc Gly	aag Lys	gag Glu	tac Tyr 120	aag Lys	tgc Cys	aag Lys	gtc Val	tcc Ser 125	aac Asn	aaa Lys	384
gcc Ala	ctc Leu	cca Pro 130	gcc Ala	ccc Pro	atc Ile	gag Glu	aaa Lys 135	acc Thr	atc Ile	tcc Ser	aaa Lys	gcc Ala 140	aaa Lys	Gly	cag Gln	432
ccc Pro	cga Arg 145	gaa Glu	cca Pro	cag Gln	gtg Val	tac Tyr 150	acc Thr	ctg Leu	ccc Pro	cca Pro	tcc Ser 155	cgg	gat Asp	gag Glu	ctg Leu	480
acc Thr 160	aag Lys	aac Asn	cag Gln	gtc Val	agc Ser 165	ctg Leu	acc Thr	tgc Cys	ctg Leu	gtc Val 170	aaa Lys	ggc	ttc	tat Tyr	ccc Pro 175	528
agc Ser	gac Asp	atc Ile	gcc Ala	gtg Val 180	gag Glu	tgg Trp	gag Glu	agc Ser	aat Asn 185	Gly	cag Gln	ccg	gag Glu	aac Asn 190	Asn	576
tac Tyr	aag Lys	acc	acg Thr	Pro	ccc Pro	gtg Val	ctg Leu	gac Asp 200	tcc Ser	gac Asp	ggc Gly	tcc Ser	Phe 205	Phe	ctc Leu	624
tac Tyr	agc Ser	Lys	Lev	acc	gtg Val	gac Asp	aag Lys 215	Ser	agg Arg	tgg Trp	cag Gln	cag Gln 220	Gly	, aac , Asn	gtc Val	672
ttc Phe	tca Ser 225	Cys	tco Ser	gto Val	g atg L Met	cat His	Glu	gct Ala	. ctg . Lev	cac His	aac Asn 235	His	tac	Thi	cag Gln	720
		· 							taa	taa	atcc	acac	7			761

Lys Ser Leu Ser Leu Ser Pro Gly Lys 240 245

<210> 1058

<211> 248

<212> PRT

<213> Artificial Sequence

<223> Description of Artificial Sequence: TNF-ALPH INHIBITOR Fc

<400> 1058

Met Asp Phe Leu Pro His Tyr Lys Asn Thr Ser Leu Gly His Arg Pro 1 5 10 15

Gly Gly Gly Gly Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala 20 25 30

Pro Glu Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro 35 40 45

Lys Asp Thr Leu Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val 50 55 60

Val Asp Val Ser His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val 65 70 75 80

Asp Gly Val Glu Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln 85 90 95

Tyr Asn Ser Thr Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln
100 105 110

Asp Trp Leu Asn Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala 115 120 125

Leu Pro Ala Pro Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro 130 135 140

Arg Glu Pro Gln Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr 145 150 155 160

Lys Asn Gln Val Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser 165 170 175

Asp Ile Ala Val Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr 180 185 190

PCT/US99/25044 WO 00/24782

Lys Thr Thr Pro Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr 200 205 195 Ser Lys Leu Thr Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe 215 Ser Cys Ser Val Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys 235 Ser Leu Ser Leu Ser Pro Gly Lys 245 <210> 1059 <211> 763 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:Fc IL-1 ANTAGONIST <220> <221> CDS <222> (4)..(747) <400> 1059 cat atg gac aaa act cac aca tgt cca cct tgt cca gct ccg gaa ctc 48 Met Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu Leu 10 5 ctg ggg gga ccg tca gtc ttc ctc ttc ccc cca aaa ccc aag gac acc Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr 25 20 ctc atg atc tcc cgg acc cct gag gtc aca tgc gtg gtg gtg gac gtg Leu Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp Val 45 40 35 age cae gaa gae cet gag gte aag tte aac tgg tae gtg gae gge gtg Ser His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly Val - 55 50

gag gtg cat aat gcc aag aca aag ccg cgg gag gag cag tac aac agc Glu Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn Ser 75 65

				gtc Val												288
				tac Tyr 100												336
				acc Thr												384
				ctg Leu												432
gtc Val	agc Ser 145	ctg Leu	acc Thr	tgc Cys	ctg Leu	gtc Val 150	aaa Lys	ggc Gly	ttc Phe	tat Tyr	ccc Pro 155	agc Ser	gac Asp	atc Ile	gcc Ala	480
gtg Val 160	gag Glu	tgg Trp	gag Glu	agc Ser	aat Asn 165	ggg Gly	cag Gln	ccg	gag Glu	aac Asn 170	aac Asn	tac Tyr	aag Lys	acc	acg Thr 175	528
cct Pro	ccc Pro	gtg Val	ctg Leu	gac Asp 180	tcc Ser	gac Asp	ggc Gly	tcc Ser	ttc Phe 185	ttc Phe	ctc Leu	tac Tyr	agc Ser	aag Lys 190	ctc Leu	576
acc Thr	gtg Val	gac Asp	aag Lys 195	agc Ser	agg <b>A</b> rg	tgg Trp	cag Gln	cag Gln 200	ggg	aac Asn	gtc Val	ttc Phe	tca Ser 205	tgc Cys	tcc Ser	624
gtg Val	atg Met	cat His 210	Glu	gct Ala	ctg Leu	cac His	aac Asn 215	His	tac Tyr	acg Thr	cag Gln	aag Lys 220	agc Ser	ctc Leu	tcc Ser	672
ctg Leu	tct Ser 225	Pro	ggt Gly	aaa Lys	ggt Gly	gga Gly 230	Gly	ggt	ggt Gly	ttc Phe	gaa Glu 235	Trp	acc Thr	ccg Pro	ggt Gly	720
tac Tyr 240	Trp	cag Gln	ccg Pro	tac Tyr	gct Ala 245	Leu	ccg Pro	ctg Leu	taa	tgga	tcc	ctcg	ag			763

<210> 1060-

<211> 248

<212> PRT

<213> Artificial Sequence

<223> Description of Artificial Sequence:Fc IL-1
ANTAGONIST

<400> 1060

Met Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu Leu Leu 1 5 10 15

- Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr Leu 20 25 30
- Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp Val Ser 35 40 45
- His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly Val Glu
  50 55 60
- Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn Ser Thr 65 70 75 80
- Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln Asp Trp Leu Asn 85 90 95
- Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu Pro Ala Pro 100 105 110
- Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro Gln 115 120 125
- Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn Gln Val 130 135 140
- Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala Val 145 150 155 160
- Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr Thr Pro 165 170 175
- Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu Thr 180 185 190
- Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser Val 195 200 205
- Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser Leu 210 215 220
- Ser Pro Gly Lys Gly Gly Gly Gly Phe Glu Trp Thr Pro Gly Tyr

225 230 235 240

Trp Gln Pro Tyr Ala Leu Pro Leu 245

<210> 1061 <211> 757

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: IL-1 ANTAGONIST Fc

<220>

<221> CDS

<222> (4)..(747)

<400> 1061

cat atg ttc gaa tgg acc ccg ggt tac tgg cag ccg tac gct ctg ccg

Met Phe Glu Trp Thr Pro Gly Tyr Trp Gln Pro Tyr Ala Leu Pro

1 5 10 15

ctg ggt gga ggc ggt ggg gac aaa act cac aca tgt cca cct tgc cca 96
Leu Gly Gly Gly Gly Asp Lys Thr His Thr Cys Pro Pro Cys Pro
20 25 30

gca cct gaa ctc ctg ggg gga ccg tca gtt ttc ctc ttc ccc cca aaa 144
Ala Pro Glu Leu Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys
35 40 45

ccc aag gac acc ctc atg atc tcc cgg acc cct gag gtc aca tgc gtg 192
Pro Lys Asp Thr Leu Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val
50 55 60

gtg gtg gac gtg agc cac gaa gac cct gag gtc aag ttc aac tgg tac

Val Val Asp Val Ser His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr

65 70 75

gtg gac ggc gtg gag gtg cat aat gcc aag aca aag ccg cgg gag gag 288
Val Asp Gly Val Glu Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu
80 85 90 95

cag tac aac agc acg tac cgt gtg gtc agc gtc ctc acc gtc ctg cac 336
Gln Tyr Asn Ser Thr Tyr Arg Val Val Ser Val Leu Thr Val Leu His
100 105 110

						ggc											384
	Gln	Asp	Trp	Leu	Asn	G1y	Lys	Glu	Tyr	Lys	Суз	Lys	Val	Ser	Asn	Lys	
				115					120					125			
	acc	ctc	cca	gcc	ccc	atc	gag	aaa	acc	atc	tcc	aaa	αcċ	aaa	aaa	cag	432
				Ala													
	WIG	neu		ALG	PIO	TTE	GIU		****			_,_	140	-3-	,		
			130					135					140				
																	400
				cca													480
	Pro	Arg	Glu	Pro	Gln	Val	Tyr	Thr	Leu	Pro	Pro	Ser	Arg	Asp	Glu	Leu	
		145					150					155					
	acc	ааσ	aac	cag	atc	agc	cta	acc	tgc	ctg	gtc	aaa	ggc	ttc	tat	CCC	528
	Thr	Lug	Agn	Gln	Val	Ser	Len	Thr	Cvs	Leu	Val	Lvs	Gly	Phe	Tyr	Pro	
		пyз	Agii	g1m	741	165			-,-		170	-	•	•	-	175	
	160					103											
		*										~~~		<b>~~</b> ~		220	576
	agc	gac	atc	gcc	gtg	gag	rgg	gag	agc	aat	999	cay	CCG	gay	2	3	2.0
	Ser	qaA	Ile	Ala	Val	Glu	Trp	Glu	Ser		GIĀ	GID	PIO	GIU		ABN	
					180					185					190		
	tac	aag	acc	acg	cct	ccc	gtg	ctg	gac	tcc	gac	ggc	tcc	ttc	ttc	ctc	624
	Tvr	Lvs	Thr	Thr	Pro	Pro	Val	Leu	Asp	Ser	Asp	Gly	Ser	Phe	Phe	Leu	
	-2-			195					200					205			
									•								•
				ctc			~~ ~	997	200	ann	taa	cag	cag	aaa	aac	atc	672
	tac	agc	aag	-	acc	grg	yac	Lag	Cor	7~~	W	Gln	Gln	Glv	lan	Va1	
	Tyr	Ser		Leu	Thr	vaı	Asp		Ser	ALY	ııp	GIII	220	013		142	
			210					215					220				
																	720
	ttc	tca	tgc	tcc	gtg	atg	cat	gag	gct	ctg	cac	aac	cac	tac	acg	cag	720
	Phe	Ser	Cys	Ser	Val	Met	His	Glu	Ala	Leu	His	Asn	His	Tyr	Thr	Gln	
		225					230					235					
	аад	ago	ctc	tcc	cta	tct	ccg	ggt	aaa	taa	tgga	tcc					757
				Ser													
	240					245		•	-								
	240					243						•					
		.0> 1															
	<21	.1> 2	48														
•	<21	.2> E	PRT														
	<21	.3> #	rtif	icia	ıl Se	quen	ce										
	<22	23> I	esci	ipti	on c	of Ar	tifi	cial	. Seq	ruenc	e:II	1 A	NTAC	ONIS	T		
			rc .	-								•					
		•	-														
	~ 4 *	10- 1	062														
	<4(	00> 1			_ m-		. 01-	, m	. M	<u>, (1</u> 1-	Dr/	יטיים כ	- A1=	Lei	r Pro	Leu	
•			s GIV	ı TT			, GT2	TAI		10		• 1.			15	5	
	1	L				5				1(	,					-	

Gly	Gly	Gly	Gly	Gly	Asp	Lys	Thr	His	Thr	Cys	Pro	Pro	Cys	Pro	Ala
			20					25					30		

- Pro Glu Leu Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro 35 40 45
- Lys Asp Thr Leu Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val 50 55 60
- Val Asp Val Ser His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val 65 70 75 80
- Asp Gly Val Glu Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln 85 90 95
- Tyr Asn Ser Thr Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln
  100 105 110
- Asp Trp Leu Asn Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala 115 120 125
- Leu Pro Ala Pro Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro 130 135 140
- Arg Glu Pro Gln Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr 145 150 155 160
- Lys Asn Gln Val Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser 165 170 175
- Asp Ile Ala Val Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr 180 185 190
- Lys Thr Thr Pro Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr 195 200 205
- Ser Lys Leu Thr Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe 210 215 220
- Ser Cys Ser Val Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys 225 230 235 240

Ser Leu Ser Leu Ser Pro Gly Lys 245

<210> 1063 <211> 773 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:Fc-VEGF ANTAGONIST <220> <221> CDS <222> (4)..(759) <400> 1063 cat atg gac aaa act cac aca tgt cca ccg tgc cca gca cct gaa ctc Met Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu Leu 10 ctg ggg gga ccg tca gtt ttc ctc ttc ccc cca aaa ccc aag gac acc Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr 25 20 ctc atg atc tcc cgg acc cct gag gtc aca tgc gtg gtg gtg gac gtg Leu Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp Val age cae gaa gae cet gag gte aag tte aac tgg tae gtg gae gge gtg 192 Ser His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly Val 60 55 50 gag gtg cat aat gcc aag aca aag ccg cgg gag gag cag tac aac agc 240 Glu Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn Ser 70 65 acg tac cgt gtg gtc agc gtc ctc acc gtc ctg cac cag gac tgg ctg 288 Thr Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln Asp Trp Leu 80 85 aat ggc aag gag tac aag tgc aag gtc tcc aac aaa gcc ctc cca gcc Asn Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu Pro Ala 105 100 ccc atc gag aaa acc atc tcc aaa gcc aaa ggg cag ccc cga gaa cca Pro Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro 115

cag gtg tac acc ctg ccc cca tcc cgg gat gag ctg acc aag aac cag

Gln Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn Gln

432

140 135 130 gtc agc ctg acc tgc ctg gtc aaa ggc ttc tat ccc agc gac atc gcc Val Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala 155 150 145 gtg gag tgg gag agc aat ggg cag ccg gag aac aac tac aag acc acg Val Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr Thr 170 160 165 cet eec gtg etg gae tee gae gge tee tte tte etc tac age aag etc 576 Pro Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu 185 180 acc gtg gac aag agc agg tgg cag cag ggg aac gtc ttc tca tgc tcc 624 Thr Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser 200 195 gtg atg cat gag gct ctg cac aac cac tac acg cag aag agc ctc tcc Val Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser 215 ctg tct ccg ggt aaa ggt ggt ggt ggt ggt gtt gaa ccg aac tgt gac. 720 Leu Ser Pro Gly Lys Gly Gly Gly Gly Val Glu Pro Asn Cys Asp 230 225 atc cat gtt atg tgg gaa tgg gaa tgt ttt gaa cgt ctg taactcgagg 769 Ile His Val Met Trp Glu Trp Glu Cys Phe Glu Arg Leu 245 240 773 atcc <210> 1064 <211> 252 <212> PRT <213> Artificial Sequence <223> Description of Artificial Sequence:Fc-VEGF ANTAGONIST <400> 1064 Met Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu Leu Leu 10

Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Asp Val Ser

. 25

Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr. Leu

30 . ....

WO 00/24782

45

His	Glu	Asp	Pro	Glu	Val	Lys	Phe	Asn	Trp	Tyr	Val	Asp	Gly	Val	Glu
	50	_				55					60	_			

40

35

Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn Ser Thr 70 75

Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln Asp Trp Leu Asn 90 85

Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu Pro Ala Pro 105

Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro Gln 120 115

Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn Gln Val 135

Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala Val 155 150 145

Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr Thr Pro 170 165

Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu Thr 185 180

Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser Val 200

Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser Leu 220 215

Ser Pro Gly Lys Gly Gly Gly Gly Val Glu Pro Asn Cys Asp Ile 235 230 225

His Val Met Trp Glu Trp Glu Cys Phe Glu Arg Leu 250 245

<210> 1065

<211> 773

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: VEGF ANTAGONIST Fc

<220>

<221> CDS

<222> (4)..(759)

<400> 1065

cat atg gtt gaa ccg aac tgt gac atc cat gtt atg tgg gaa tgg gaa 49
Met Val Glu Pro Asn Cys Asp Ile His Val Met Trp Glu Trp Glu

1 5 10 15

tgt ttt gaa cgt ctg ggt ggt ggt ggt ggt gac aaa act cac aca tgt 96
Cys Phe Glu Arg Leu Gly Gly Gly Gly Asp Lys Thr His Thr Cys
20 25 30

cca ccg tgc cca gca cct gaa ctc ctg ggg gga ccg tca gtt ttc ctc 144
Pro Pro Cys Pro Ala Pro Glu Leu Leu Gly Gly Pro Ser Val Phe Leu
35 40 45

ttc ccc cca aaa ccc aag gac acc ctc atg atc tcc cgg acc cct gag

Phe Pro Pro Lys Pro Lys Asp Thr Leu Met Ile Ser Arg Thr Pro Glu

50 55 60

gtc aca tgc gtg gtg gtg gac gtg agc cac gaa gac cct gag gtc aag 240
Val Thr Cys Val Val Val Asp Val Ser His Glu Asp Pro Glu Val Lys
65 70 75

ttc aac tgg tac gtg gac ggc gtg gag gtg cat aat gcc aag aca aag 288

Phe Asn Trp Tyr Val Asp Gly Val Glu Val His Asn Ala Lys Thr Lys

80 85 90 95

ccg cgg gag gag cag tac aac agc acg tac cgt gtg gtc agc gtc ctc 336
Pro Arg Glu Glu Gln Tyr Asn Ser Thr Tyr Arg Val Val Ser Val Leu
100 105 110

acc gtc ctg cac cag gac tgg ctg aat ggc aag gag tac aag tgc aag
Thr Val Leu His Gln Asp Trp Leu Asn Gly Lys Glu Tyr Lys Cys Lys

115
120
125

gtc tcc aac aaa gcc ctc cca gcc ccc atc gag aaa acc atc tcc aaa 432
Val Ser Asn Lys Ala Leu Pro Ala Pro Ile Glu Lys Thr Ile Ser Lys
130 135 140

gcc aaa ggg cag ccc cga gaa cca cag gtg tac acc ctg ccc cca tcc 480
Ala Lys Gly Gln Pro Arg Glu Pro Gln Val Tyr Thr Leu Pro Pro Ser

145 150 155

				acc Thr												528
				agc Ser 180												576
				tac Tyr												624
				tac Tyr												672
cag Gln	ggg Gly 225	aac Asn	gtc Val	ttc Phe	tca Ser	tgc Cys 230	tcc Ser	gtg Val	atg Met	cat His	gag Glu 235	gct Ala	ctg Leu	cac His	aac Asn	720
				aag Lys									taa	ctcg	agg	769
atc	3															773
<21: <21: <21:		52 RT rtif escr		l Se	-		cial	Seq	uenc	e:VE	GF A	NTAG	onis	T	•	
	0> 1 Val		Pro	Asn 5		Asp	Ile	His	Val		Trp	Glu	Trp	Glu 15	Сув	
Phe	Glu	. Arg	Leu 20		Gly	Gly	Gly		Asp	Lys	Thr	His	Thr 30		Pro	
Pro	Cys	Pro		Pro	Glu	Leu	Leu 40		'Gly	Pro	Ser	Val		e Lev	Phe	
Pro	Pro		Pro	Lys	a Asy	Thr 55		Met	: Ile	e Ser	Arg	Thr	Pro	G1	val	

Thr Cys Val Val Val Asp Val Ser His Glu Asp Pro Glu Val Lys Phe 65 70 75 80

- Asn Trp Tyr Val Asp Gly Val Glu Val His Asn Ala Lys Thr Lys Pro 85 90 95
- Arg Glu Glu Gln Tyr Asn Ser Thr Tyr Arg Val Val Ser Val Leu Thr 100 105 110
- Val Leu His Gln Asp Trp Leu Asn Gly Lys Glu Tyr Lys Cys Lys Val 115 120 125
- Ser Asn Lys Ala Leu Pro Ala Pro Ile Glu Lys Thr Ile Ser Lys Ala 130 135 140
- Lys Gly Gln Pro Arg Glu Pro Gln Val Tyr Thr Leu Pro Pro Ser Arg 145 150 155 160
- Asp Glu Leu Thr Lys Asn Gln Val Ser Leu Thr Cys Leu Val Lys Gly
  165 170 175
- Phe Tyr Pro Ser Asp Ile Ala Val Glu Trp Glu Ser Asn Gly Gln Pro 180 185 190
- Glu Asn Asn Tyr Lys Thr Thr Pro Pro Val Leu Asp Ser Asp Gly Ser 195 200 205
- Phe Phe Leu Tyr Ser Lys Leu Thr Val Asp Lys Ser Arg Trp Gln Gln 210 215 220
- Gly Asn Val Phe Ser Cys Ser Val Met His Glu Ala Leu His Asn His 225 230 235 240
- Tyr Thr Gln Lys Ser Leu Ser Leu Ser Pro. Gly Lys 245 250

<210> 1067

<211> 748

<212> DNA

<213> Artificial Sequence

<220>

<220>

<221> CDS <222> (4)..(732)

160

<400	> 10	67														
cat	atg	gac	aaa	act	cac	aca	tgt	cca	cct	tgt	cca	gct	ccg	gaa	ctc	48
	Met	Asp	Lys	Thr	His	Thr	Cys	Pro	Pro	Сув	Pro	Ala	Pro	Glu		
	1				5					10					15	
ctg	ggg	gga	ccg	tca	gtc	ttc	ctc	ttc	CCC	cca	aaa	CCC	aag	gac	āÇC	96
Leu	Gly	Gly	Pro	Ser	Val	Phe	Leu	Phe	Pro	Pro	Lys	Pro	Lys	Asp	Thr	
				20					25					30		
ctc	atg	atc	tcc	cgg	acc	cct	gag	gtc	aca	tgc	gtg	gtg	gtg	gac	gtg	144
Leu	Met	Ile	Ser	Arg	Thr	Pro	Glu	Val	Thr	Cys	Val	Val	Val	Asp	Val	
			35					40					45			
agc	cac	gaa	gac	cct	gag	gtc	aag	ttc	aac	tgg	tac	gtg	gac	ggc	gtg	192
Ser	His	Glu	Asp	Pro	Glu	Val	Lys	Phe	Asn	Trp	Tyr	Val	Asp	Gly	Val	
		50	_				55					60				
gag	ata	cat	aat	gcc	aag	aca	aag	ccg	cgg	gag	gag	cag	tac	aac	agc	240
Glu	Val	His	Asn	Ala	Lys	Thr	Lys	Pro	Arg	Glu	Glu	Gln	Tyr	Asn	Ser	
	65				_	70					75					•
	•															•
асσ	tac	cgt	ata	atc	agc	gtc	ctc	acc	gtc	ctg	cac	cag	gac	tgg	ctg	288
Thr	Tvr	Arg	Val	Val	Ser	Val	Leu	Thr	Val	Leu	His	Gln	Asp	Trp	Leu	
80					85					90					95	
																•
aat	aac	aád	dad	tac	aag	tgc	aag	gtc	tcc	aac	aaa	gcc	ctc	cca	gcc	336
Asn	Glv	Lvs	Glu	Tyr	Lys	Cys	Lys	Val	Ser	Asn	Lys	Ala	Leu	Pro	Ala	
				100		_			105					110		
ccc	ato	дад	aaa	acc	atc	tcc	aaa	gcc	aaa	ggg	cag	ccc	cga	gaa	cca	384
Pro	Ile	Glu	Lvs	Thr	Ile	Ser	Lys	Ala	Lys	Gly	Gln	Pro	Arg	Glu	Pro	
			115					120					125	i		
cad	ata	tac	acc	cto	ccc	: cca	tcc	cgg	gat	gag	ctg	acc	: aag	aac	cag	432
Gln	Val	Tvr	Thr	Leu	Pro	Pro	Ser	Arg	Asp	Glu	Leu	Thr	Lys	Asr	Gln	
		130					135	i				140	)			
									•							
ato	. 200	. cto	acc	: tac	: cto	ato	aaa	ggo	tto	: tat	ccc	ago	gad	ato	gcc	480
Val	Set	Len	The	Cvs	Lev	Val	Lys	G13	Phe	Tyr	Pro	Ser	: Ası	Ile	a Ala	
	145					150					155	5				
	47.	•														
-	7 (72/	ı tar	, dai	1 200	aat	. aac	cac	cc	g gaç	aac	aac	: tac	aaq	g acc	c acg	52
17:51	, yay	بيشى ، بريم ه	, yu.	, ~y`	r Agr	1 G1:	, Gli	r Pro	G11	ı Asr	a Asi	ı Ty	r Ly	s Th	r Thir	
160			, 51	. 56	165			, _ ,		170	)				175	
TO	3					-										

cct ccc gtg ctg gac tcc gac ggc tcc ttc ttc ctc tac agc aag ctc Pro Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu 185 180 acc gtg gac aag agc agg tgg cag cag ggg aac gtc ttc tca tgc tcc Thr Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser 200 195 gtg atg cat gag gct ctg cac aac cac tac acg cag aag agc ctc tcc 672 Val Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser 210 215 ctg tct ccg ggt aaa ggt gga ggt ggt tgc acc acc cac tgg ggt 720 Leu Ser Pro Gly Lys Gly Gly Gly Gly Cys Thr Thr His Trp Gly 235 230 225 748 ttc acc ctg tgc taatggatcc ctcgag Phe Thr Leu Cys 240 <210> 1068 <211> 243 <212> PRT <213> Artificial Sequence <223> Description of Artificial Sequence:Fc-MMP INHIBITOR <400> 1068 Met Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu Leu Leu 5 Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr Leu 30 25 20 Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp Val Ser 35 40 His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly Val Glu - 60 55 Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn Ser Thr 70 65 Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln Asp Trp Leu Asn . 90 Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu Pro Ala Pro

100 105 110

Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro Gln
115 120 125

Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn Gln Val 130 135 140

Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala Val 145 150 155 160

Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr Thr Pro 165 170 175

Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu Thr 180 185 190

Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser Val 195 200 205

Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser Leu 210 215 220

Ser Pro Gly Lys Gly Gly Gly Gly Gly Cys Thr Thr His Trp Gly Phe 225 230 235 240

Thr Leu Cys

<210> 1069

<211> 763

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: MMP INHIBITOR Fc

<220>

<221> CDS

<222> (4)..(753)

<400> 1069

cat atg tgc acc acc cac tgg ggt ttc acc ctg tgc ggt gga ggc ggt

Met Cys Thr Thr His Trp Gly Phe Thr Leu Cys Gly Gly Gly

1 10 15

Gly	gac Asp	aaa Lys	ggt Gly	gga Gly 20	ggc Gly	ggt Gly	Gly	gac Asp	aaa Lys 25	act Thr	cac His	aca Thr	tgt Cys	cca Pro 30	cct Pro	96
											gtt Val					144
											acc Thr					192
tgc Cys	gtg Val 65	gtg. Val	gtg Val	gac Asp	gtg Val	agc Ser 70	cac His	gaa Glu	gac Asp	cct Pro	gag Glu 75	gtc Val	aag Lys	ttc Phe	aac Asn	240
tgg Trp 80	tac Tyr	gtg Val	gac Asp	ggc Gly	gtg Val 85	gag Glu	gtg Val	cat His	aat Asn	gcc Ala 90	aag Lys	aca Thr	aag Lys	ccg Pro	cgg Arg 95	288
gag Glu	gag Glu	cag Gln	tac Tyr	aac Asn 100	agc Ser	acg Thr	tac Tyr	cgt Arg	gtg Val 105	gtc Val	agc Ser	gtc Val	ctc Leu	acc Thr 110	gtc Val	336
ctg Leu	cac His	cag Gln	gac Asp 115	tgg Trp	ctg Leu	aat Asn	ggc	aag Lys 120	gag Glu	tac Tyr	aag Lys	tgc Cys	aag Lys 125	gtc Val	tcc Ser	384
aac Asn	aaa Lys	gcc Ala 130	ctc Leu	cca Pro	gcc Ala	ccc	atc Ile 135	Glu	aaa Lys	acc Thr	atc Ile	tcc Ser 140	aaa Lys	gcc Ala	aaa Lys	432
ggg Gly	cag Gln 145	Pro	cga Arg	gaa Glu	cca Pro	cag Gln 150	Val	tac Tyr	acc	ctg Leu	ccc Pro 155	Pro	tcc	cgg	gat	480
gag Glu 160	Leu	acc Thr	aag	aac	cag Gln 165	Val	ago Ser	ctg Leu	acc Thr	tgo Cys	Leu	gtc Val	aaa Lys	ggc Gly	Phe	528
tat Tyr	ccc	ago Ser	gac Asp	ato	Ala	gtç Val	gaq Glu	tgg Trp	gag Glu 185	ı Ser	aat Asn	ggg Gly	caç Glr	p ccq 1 Pro 190	gag Glu	576
aac Asn	aac Asr	tac Tyr	aag Lys	Thr	acç Thr	cct Pro	cco Pro	gtq Val	Let	g gad	tco Ser	gac Asp	gg( G1;	y 56.	ttc Phe	624

ttc ctc tac agc aag ctc acc gtg gac aag agc agg tgg cag cag ggg Phe Leu Tyr Ser Lys Leu Thr Val Asp Lys Ser Arg Trp Gln Gln Gly 215 210 aac gtc ttc tca tgc tcc gtg atg cat gag gct ctg cac aac cac tac 720 Asn Val Phe Ser Cys Ser Val Met His Glu Ala Leu His Asn His Tyr 230 225 763 acg cag aag agc ctc tcc ctg tct ccg ggt aaa taatggatcc Thr Gln Lys Ser Leu Ser Leu Ser Pro Gly Lys 245 240 <210> 1070 <211> 250 <212> PRT <213> Artificial Sequence <223> Description of Artificial Sequence: MMP INHIBITOR FC <400> 1070 Met Cys Thr Thr His Trp Gly Phe Thr Leu Cys Gly Gly Gly Gly 5 1 Asp Lys Gly Gly Gly Gly Asp Lys Thr His Thr Cys Pro Pro Cys 25 20 Pro Ala Pro Glu Leu Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro 40 Lys Pro Lys Asp Thr Leu Met Ile Ser Arg Thr Pro Glu Val Thr Cys 55 Val Val Val Asp Val Ser His Glu Asp Pro Glu Val Lys Phe Asn Trp 70. 65 Tyr Val Asp Gly Val Glu Val His Asn Ala Lys Thr Lys Pro Arg Glu 85 Glu Gln Tyr Asn Ser Thr Tyr Arg Val Val Ser Val Leu Thr Val Leu 105 100 His Gln Asp Trp Leu Asn Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn 120 115 Lys Ala Leu Pro Ala Pro Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly 140 135

Gln Pro Arg Glu Pro Gln Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu 145 150 155 160

Leu Thr Lys Asn Gln Val Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr 165 170 175

Pro Ser Asp Ile Ala Val Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn 180 185 190

Asn Tyr Lys Thr Thr Pro Pro Val Leu Asp Ser Asp Gly Ser Phe Phe 195 200 205

Leu Tyr Ser Lys Leu Thr Val Asp Lys Ser Arg Trp Gln Gln Gly Asn 210 215 220

Val Phe Ser Cys Ser Val Met His Glu Ala Leu His Asn His Tyr Thr 225 230 235 240

Gln Lys Ser Leu Ser Leu Ser Pro Gly Lys 245 250

<210> 1071

<211> 13

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: INTEGRIN BINDING PEPTIDE

<400> 1071

Cys Gly Arg Glu Cys Pro Arg Leu Cys Gln Ser Ser Cys
1 5 10

<210> 1072

<211> 13

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: INTEGRIN BINDING PEPTIDE

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<400> 1072
Cys Asn Gly Arg Cys Val Ser Gly Cys Ala Gly Arg Cys
                5
<210> 1073
<211> 8
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: INTEGRIN
      BINDING PEPTIDE
<400> 1073
Cys Leu Ser Gly Ser Leu Ser Cys
                 5
 1
<210> 1074
<211> 6
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: INTEGRIN
      BINDING PEPTIDE
<400> 1074
Asn Gly Arg Ala His Ala
 <210> 1075
 <211> 5
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: INTEGRIN
     BINDING PEPTIDE
 <220>
 <221> CDS
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<222> (10)..(189)

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<400> 1075
Cys Asn Gly Arg Cys
<210> 1076
<211> 9
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: INTEGRIN
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<400> 1076
Cys Asp Cys Arg Gly Asp Cys Phe Cys
                5
<210> 1077
<211> 7
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: INTEGRIN
      BINDING PEPTIDE
<400> 1077
Cys Gly Ser Leu Val Arg Cys
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<210> 1078
<211> 8
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: INTEGRIN
      BINDING PEPTIDE
<400> 1078 .
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Arg Thr Asp Leu Asp Ser Leu Arg

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5 1

<210> 1079

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: INTEGRIN BINDING PEPTIDE

<400> 1079 .

Gly Asp Leu Asp Leu Leu Lys Leu Arg Leu Thr Leu 5 .

<210> 1080

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: INTEGRIN-BINDING PEPTIDE

<400> 1080

Gly Asp Leu His Ser Leu Arg Gln Leu Leu Ser Arg 5 1

<210> 1081

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: INTEGRIN-BINDING PEPTIDE

<400> 1081

Arg Asp Asp Leu His Met Leu Arg Leu Gln Leu Trp 1 ... 5 . 10

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<210> 1082
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial
      Sequence: INTEGRIN-BINDING PEPTIDE
<400> 1082
Ser Ser Asp Leu His Ala Leu Lys Lys Arg Tyr Gly
                                    10
                 5
<210> 1083
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial
      Sequence: INTEGRIN-BINDING PEPTIDE
<400> 1083
Arg Gly Asp Leu Lys Gln Leu Ser Glu Leu Thr Trp
 1 . 5
<210> 1084
<211> 12
<212> PRT
<213> Artificial Sequence
 <220>
 <223> Description of Artificial
       Sequence: INTEGRIN-BINDING PEPTIDE
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 Arg Gly Asp Leu Ala Ala Leu Ser Ala Pro Pro Val
                                     10
                   5
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<210> 1085 <211> 15

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<212> PRT
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<213> Artificial Sequence

<220>

<400> 1085

Asp Phe Leu Pro His Tyr Lys Asn Thr Ser Leu Gly His Arg Pro 1 5 10 15

<210> 1086

<211> 18

<212> PRT

<213> Artificial Sequence

<220>

<400> 1086

Gly Glu Arg Trp Cys Phe Asp Gly Pro Leu Thr Trp Val Cys Gly Glu
1 5 10 15

Glu Ser

<210> 1087

<211> 20

<212> PRT

<213> Artificial Sequence

<220>

<400> 1087

Arg Gly Trp Val Glu Ile Cys Val Ala Asp Asp Asn Gly Met Cys Val 1 5 10 15

Thr Glu Ala Gln

... 20

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<210> 1088
<211> 19
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VEGF ANTAGONIST
     PEPTIDE
<400> 1088
Gly Trp Asp Glu Cys Asp Val Ala Arg Met Trp Glu Trp Glu Cys Phe
                                  10
                5
Ala Gly Val
<210> 1089
<211> 16
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: VEGF ANTAGONIST
     PEPTIDE
<400> 1089
Arg Gly Trp Val Glu Ile Cys Glu Ser Asp Val Trp Gly Arg Cys Leu
                                                      15
                            10
               5
 1
<210> 1090
<211> 16
<212> PRT
<213> Artificial Sequence
 <220>
<223> Description of Artificial Sequence: VEGF ANTAGONIST
      PEPTIDE
 <400> 1090
 Arg Gly Trp Val Glu Ile Cys Glu Ser Asp Val Trp Gly Arg Cys Leu
                                                   . 15
  1 ... 5
                       . 10
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<210> 1091
<211> 19
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: VEGF ANTAGONIST
      PEPTIDE
<400> 1091
Gly Gly Asn Glu Cys Asp Ile Ala Arg Met Trp Glu Trp Glu Cys Phe
                5
Glu Arg Leu
<210> 1092
<211> 16
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: VEGF ANTAGONIST
      PEPTIDE
<400> 1092
Arg Gly Trp Val Glu Ile Cys Ala Ala Asp Asp Tyr Gly Arg Cys Leu
                                    10
                  5
<210> 1093
<211> 8
 <212> PRT
 <213> Artificial Sequence
<220>
 <223> Description of Artificial Sequence: MMP INHIBITOR
       PEPTIDE
 <400> 1093
 Cys Leu Arg Ser Gly Xaa Gly Cys
       ... 5
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<210> 1094
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:MMP INHIBITOR
      PEPTIDE
<400> 1094
Cys Xaa Xaa His Trp Gly Phe Xaa Xaa Cys
                5
<210> 1095
<211> 5
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: MMP INHIBITOR
      PEPTIDE
<400> 1095
Cys Xaa Pro Xaa Cys
<210> 1096
<211> 10
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence: MMP INHIBITOR
      PEPTIDE
 <400> 1096
 Cys Arg Arg His Trp Gly Phe Glu Phe Cys
           5
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<210> 1097 <211> 10

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<212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: MMP INHIBITOR
      PEPTIDE
 <400> 1097
 Ser Thr Thr His Trp Gly Phe Thr Leu Ser
 1 5
 <210> 1098
 <211> 10
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:MMP INHIBITOR
      PEPTIDE
 <400> 1098
 Cys Ser Leu His Trp Gly Phe Trp Trp Cys
  1 5 10
 <210> 1099
 <211> 15
<212> PRT
 <213> Artificial Sequence
 <223> Description of Artificial Sequence: CARBOHYDRATE
       (GD1 ALPHA) MIMETIC PEPTIDE
 <400> 1099
 Trp His Trp Arg His Arg Ile Pro Leu Gln Leu Ala Ala Gly Arg
                                                    15
                                  10
                  5
 <210> 1100
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<211> 6 <212> PRT ...

<213> Artificial Sequence

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<220>
 <223> Description of Artificial Sequence:BETA-2 GP1AB
      BINDING PEPTIDE
 <400> 1100
 Leu Lys Thr Pro Arg Val
                  5
  1
 <210> 1101
 <211> 8
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence:BETA-2 GP1AB
      BINDING PEPTIDE
 <400> 1101
 Asn Thr Leu Lys Thr Pro Arg Val
                 5
 <210> 1102
 <211> 11
 <212> PRT
 <213> Artificial Sequence
<220>
 <223> Description of Artificial Sequence:BETA-2 GP1AB
      BINDING PROTEIN
 <400> 1102
 Asn Thr Leu Lys Thr Pro Arg Val Gly Gly Cys
                   5
   1
 <210> 1103
 <211> 6
 <212> PRT
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<223> Description of Artificial Sequence:BETA-2 GP1AB

<213> Artificial Sequence

BINDING PROTEIN

<220> ...

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<400> 1103
Lys Asp Lys Ala Thr Phe
1 5
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<210> 1104

<211> 10

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:BETA-1 GP1AB BINDING PROTEIN

<400> 1104

Lys Asp Lys Ala Thr Phe Gly Cys His Asp 1 5 10

<210> 1105

<211> 12

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:BETA-2 GP1AB BINDING PEPTIDE

<400> 1105

Lys Asp Lys Ala Thr Phe Gly Cys His Asp Gly Cys . 1 5 10

<210> 1106

<211> 6

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:BETA-2 GP1AB BINDING PROTEIN

<400> 1106

Thr Leu Arg Val Tyr Lys

1 5

<210> 1107

<211> 9

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:BETA-2 GP1AB BINDING PROTEIN

<400> 1107

Ala Thr Leu Arg Val Tyr Lys Gly Gly

1 5

<210> 1108

<211> 10

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:BETA-2 GP1AB BINDING PROTEIN

<400> 1108

Cys Ala Thr Leu Arg Val Tyr Lys Gly Gly
1 5 10

<210> 1109

<211> 14

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:MEMBRANE
 TRANSPORTING PEPTIDE

<400> 1109

Ile Asn Leu Lys Ala Leu Ala Ala Leu Ala Lys Lys Ile Leu

... 5 . 10

```
<210> 1110
 <211> 12
 <212> PRT
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 <223> Description of Artificial Sequence: MEMBRANE
       TRANSPORTING PEPTIDE
 <400> 1110
 Gly Trp Thr Leu Asn Ser Ala Gly Tyr Leu Leu Gly
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 <210> 1111
 <211> 27
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence: MEMBRANE
       TRANSPORTING PEPTIDE
 <400> 1111
 Gly Trp Thr Leu Asn Ser Ala Gly Tyr Leu Leu Gly Lys Ile Asn Leu
 Lys Ala Leu Ala Leu Ala Lys Lys Ile Leu
              20
 <210> 1112
<211> 22
 <212> DNA
 <213> Artificial Sequence
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 <223> Description of Artificial Sequence:FC PCR PRIMER
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                                                                   22
aacataagta cctgtaggat cg
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<210> 1113 <211> 81

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<212> DNA
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<223> Description of Artificial Sequence:Fc-TNF ALPHA
      PCR PRIMER
<220>
<221> CDS
<222> (1)..(126)
<400> 1113
ccg cgg atc cat tac gga cgg tga ccc aga gag gtg ttt ttg tag tgc 48
Pro Arg Ile His Tyr Gly Arg Pro Arg Glu Val Phe Leu Cys
                 5
                                                                 81
ggc agg aag tca cca cct cca cct tta ccc
Gly Arg Lys Ser Pro Pro Pro Pro Pro Leu Pro
            20
<210> 1114
<211> 7
<212> PRT
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<223> Description of Artificial Sequence:Fc-TNF ALPHA
      PCR PRIMER
<400> 1114
Pro Arg Ile His Tyr Gly Arg
                5
1
<210> 1115
<211> 6
<212> PRT
<213> Artificial Sequence
<223> Description of Artificial Sequence:Fc-TNF ALPHA
      PCR PRIMER
<400> 1115
Pro Arg Glu Val Phe Leu
<210> 1116 ___
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<211> 12 <212> PRT

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<213> Artificial Sequence
<223> Description of Artificial Sequence:Fc-TNF ALPHA
     PCR PRIMER
<400> 1116
Cys Gly Arg Lys Ser Pro Pro Pro Pro Pro Leu Pro
                5
<210> 1117
<211> 81
<212> DNA
<213> Artificial Sequence
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<223> Description of Artificial Sequence:TNF-ALPHA
      INHIBITOR-FC PCR PRIMER
<400> 1117
gaataacata tggacttcct gccgcactac aaaaacacct ctctgggtca ccgtccgggt 60
ggaggcggtg gggacaaaac t
<210> 1118
<211> 81
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      PCR PRIMER
<400> 1118
ccgcggatcc attacagcgg cagagcgtac ggctgccagt aacccggggt ccattcgaaa 60
ccaccacctc cacctttacc c
<210> 1119
<211> 81
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: IL-1 ANTAGONIST
      -Fc PCR PRIMER
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<400> 1119

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gaataacata tgttcgaatg gaccccgggt tactggcagc cgtacgctct gccgctgggt 60
ggaggcggtg gggacaaaac t
<210> 1120
<211> 57
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:Fc-VEGF
      ANTAGONIST OLIGONUCLEOTIDE
<400> 1120
gttgaaccga actgtgacat ccatgttatg tgggaatggg aatgttttga acgtctg 57
<210> 1121
<211> 57
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:Fc-VEGF
      ANTAGONIST OLIGONUCLEOTIDE
<400> 1121
cagacgttca aaacattccc attcccacat aacatggatg tcacagttcg gttcaac 57
<210> 1122
<211> 57
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:Fc-VEGF
      ANTAGONIST PCR TEMPLATE
<400> 1122
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                                                                  57
<210> 1123
<211> 48
<212> DNA ...
<213> Artificial Sequence
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PCT/US99/25044 WO 00/24782 <220> <223> Description of Artificial Sequence:Fc PRIMER <400> 1123 48 atttgattct agaaggagga ataacatatg gacaaaactc acacatgt <210> 1124 <211> 51 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:Fc PRIMER <400> 1124 gtcacagttc ggttcaacac caccaccacc acctttaccc ggagacaggg a 51 <210> 1125 <211> 54 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:Fc-VEGF ANTAGONIST PCR PRIMER <400> 1125 tecetytete egggtaaagg tggtggtggt ggtgttgaac egaactgtga cate 54 <210> 1126 <211> 39 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence: Fc-VEGF ANTAGONIST-FC PCR PRIMER <400> 1126 39

<210> 1127---<211> 48

<212> DNA

ccgcggatcc tcgagttaca gacgttcaaa acattccca

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<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence:Fc-VEGF
      ANTAGONIST-FC PCR PRIMER
<400> 1127
                                                                  48
atttgattct agaaggagga ataacatatg gttgaaccga actgtgac
<210> 1128
<211> 51
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Ala Ala Arg Ala
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## (19) World Intellectual Property Organization International Bureau



## - 1 Marie (1940) ( 1 April 1 A

### (43) International Publication Date 4 May 2000 (04.05.2000)

(51) International Patent Classification7:

#### **PCT**

# (10) International Publication Number WO 00/24782 A3

- C12N 15/62, 15/70, 1/21
- (21) International Application Number: PCT/US99/25044
- (22) International Filing Date: 25 October 1999 (25.10.1999)
- (25) Filing Language:

English

C07K 19/00,

(26) Publication Language:

English

(30) Priority Data:

60/105.371 09/428,082 23 October 1998 (23.10.1998) US 22 October 1999 (22.10.1999) US

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- (81) Designated States (national): AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW). Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

#### Published:

- with international search report
- (88) Date of publication of the international search report: 6 June 2002

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

A3

#### (54) Title: MODIFIED PEPTIDES, COMPRISING AN FC DOMAIN, AS THERAPEUTIC AGENTS

(57) Abstract: The present invention concerns fusion of Fc domains with biologically active peptides and a process for preparing pharmaceutical agents using biologically active peptides. In this invention, pharmacologically active compounds are prepared by a process comprising: a) selecting at least one peptide that modulates the activity of a protein of interest; and b) preparing a pharmacologic agent comprising an Fc domain covalently linked to at least one amino acid of the selected peptide. Linkage to the vehicle increases the half-life of the peptide, which otherwise would be quickly degraded *in vivo*. The preferred vehicle is an Fc domain. The peptide is preferably selected by phage display, *E. coli* display, ribosome display, RNA-peptide screening, or chemical-peptide screening.

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A. CLASSIFICATION OF SUBJECT MATTER IPC 7 CO7K19/00 C12N15/62 C12N15/70 C12N1/21 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 7 C07K A61K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) BIOSIS, EMBASE, WPI Data, PAJ, EPO-Internal, STRAND C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. X WO 98 46257 A (AMGEN INC.) 1-3,5-722 October 1998 (1998-10-22) page 3, line 12 -page 4, line 4 page 12, line 9 - line 25 Y 11-21,51X WO 96 18412 A (BETH ISRAEL HOSPITAL 1-3,5,6, ASSOCIATION) 20 June 1996 (1996-06-20) 22-24 page 8, line 14 -page 12, line 26 claims X Further documents are listed in the continuation of box C. Patent family members are listed in annex. Special categories of cited documents : "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the *A* document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) involve an inventive step when the document is taken alone document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 07 12 2000 18 October 2000 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Ruswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3016 Nooij, F

Int .tional Application No PCT/US 99/25044

C./Continue	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	FC1703 99/25044
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tional application No.
PCT/US 99/25044

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This Inte	emational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1.	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2.	Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
з. 🗌	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Inte	ernational Searching Authority found multiple inventions in this international application, as follows:
	see additional sheet
	As a result of the prior review under R. 40.2(e) PCT, no additional fees are to be refunded.
1. X	As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3	As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. 🗌	No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remari	k on Protest  X The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.

## FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1-7 (partially), 8-11 (completely), 22-32 (partially), 35 (completely), 39-51 (partially)

Compositions of matter of the formula (X1)a-F1-(X2)b and multimers thereof, wherein F1 is an Fc domain, X1 and X2 are each independently selected from -(L1)c-P1, -(L1)c-P1-(L2)d-P2, -(L1)c-P1-(L2)d-P2-(L3)e-P3, and -(L1)c-P1-(L2)d-P2-(L3)e-P3-(L4)f-P4. P1, P2, P3 and P4 are each independently sequences of pharmacologically activbe peptides; L1, L2, L3 and L4 are each independently linkers, and a, b, c, d and e are each independently 0 or 1, provided that at least one of a and b is 1; DNA encoding said composition, an expression vector comprising said DNA, a host cell comprising said expression vector, Proces for preparing a pharmacologically active compound, and wherein X1 and X2 comprise an IL-1 antagonist peptide sequence.

2. Claims: 1-7 (partially), 12-17 (completely), 22-32 (partially), 33 (completely), 39-51 (partially)

As in subject 1, but wherein X1 and X2 comprise an EPO-mimetic peptide sequence.

 Claims: 1-7 (partially), 18-21 (completely), 22-32 (partially), 34 (completely), 39-51 (partially)

As in subject 1, but wherein P1 is a TP0-mimetic peptide sequence  $\ \ \,$ 

4. Claims: 26-32 (partially), 36 (completely), 39-51 (partially)

Process for preparing a pharmacologically active compound, which comprises selecting at least one randomized peptide that modulates the activity of a protein of interest, and preparing a pharmacologic agent comprising one Fc domain covalently linked to at least one amino acid sequence of the selected peptide(s); wherein said peptide is an MMP inhibitor peptide or a VEGF antagonist peptide.

5. Claims: 26-32 (partially), 37 (completely),

## FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

39-51 (partially)

6. Claims: 26-32 (partially), 38 (completely), 39-51 (partially)

As in subject 4, but wherein said peptide is a CTLA4 mimetic peptide.

ormation on patent family members

Interretonal Application No
PC 1, US 99/25044

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